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<b>(21) International Application Number:</b> PCT/AU89/00123 <b>(22) International Filing Date:</b> 23 March 1989 (23.03.89)  <b>(31) Priority Application Number:</b> PI 7391 <b>(32) Priority Date:</b> 23 March 1988 (23.03.88) <b>(33) Priority Country:</b> AU  <b>(71) Applicant (for all designated States except US):</b> THE UNIVERSITY OF MELBOURNE [AU/AU]; Parkville, VIC 3052 (AU).  <b>(72) Inventors; and</b> <b>(75) Inventors/Applicants (for US only) :</b> SINGH, Mohan, Bir [IN/AU]; 7 Lloyd Court, Templestowe, VIC 3106 (AU). HOUGH, Terryn [AU/AU]; 25 Bowman Street, Mordialloc, VIC 3195 (AU). THEERAKULPISUT, Piyada [TH/AU]; 1/74 Canning Street, Carlton, VIC 3053 (AU). KNOX, Robert, Bruce [AU/AU]; 274 Balwyn Road, North Balwyn, VIC 3104 (AU).		<b>(74) Agents:</b> NOONAN, Gregory, J. et al.; Davies & Collison, 1 Little Collins Street, Melbourne, VIC 3000 (AU).  <b>(81) Designated States:</b> AT (European patent), BE (European patent), CH (European patent), DE (European patent), DK, FR (European patent), GB (European patent), IT (European patent), JP, LU (European patent), NL (European patent), SE (European patent), US.  <b>Published</b> <i>With international search report.</i>
<b>(54) Title:</b> RYEGRASS POLLEN ALLERGEN  <b>(57) Abstract</b>  <p>The major allergenic protein <i>LOI pI</i> from pollen of ryegrass <i>Lolium perenne</i>L. is produced by recombinant DNA techniques. The DNA sequence encoding the above protein, expression vectors, host transformed and cell lines containing the coding sequence for <i>LOI pI</i> protein are also described. The use of the above DNA sequences and recombinant protein in nucleic hybridization, tissue specificity diagnosis and detection of specific antibodies in biological samples are also disclosed. The possible use of the promoter sequence of <i>LOI pI</i> in the developmental regulation of <i>LOI pI</i> gene expression or any other gene during the development of the pollen, in inhibiting pollen development or function and inducing nuclear male sterility are also disclosed.</p>		

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1                   "RYEGRASS POLLEN ALLERGEN"

3           The present invention relates to the major  
allergenic protein Lol pI from pollen of ryegrass,  
5 Lolium perenne L. and to derivatives and homologues  
thereof and to allergenic proteins immunologically related  
7 thereto. More particularly, the present invention is  
directed to recombinant Lol pI, and its derivatives, and  
9 to an expression vector capable of directing synthesis of  
same. Even more particularly, the present invention is  
11 directed to cDNA encoding Lol pI and to its promoter and  
to an expression vector comprising same.

13

          Allergens constitute the most abundant proteins of  
15 grass pollen, which is the major cause of allergic disease  
in temperate climates (MARSH, 1975, HILL et al., 1979).  
17 The first descriptions of their allergenic proteins showed  
that they are immunochemically distinct, and are known as  
19 groups I, II, III and IV (JOHNSON AND MARSH 1965, 1966).  
Using the recently proposed International Union of  
21 Immunological Societies' (IUIS) nomenclature, these  
allergens are designated Lol pI, Lol pII, Lol pIII and  
23 Lol pIV. The major allergen, Lol pI is an acidic  
glycoprotein of molecular weight ca.32 kD and comprises  
25 four isoallergenic variants. The other minor allergens  
isolated from ryegrass pollen range in molecular weight  
27 from 10 to 76 kD (see review by FORD AND BALDO, 1986). The  
allergen Lol pI constitutes ca.5% of the total extracted  
29 pollen proteins and is a glycoprotein (HOWLETT & CLARKE,  
1981) containing a 5% carbohydrate moiety. Studies with  
31 carbohydrate splitting have demonstrated that the  
carbohydrate does not contribute to the allergic  
33 response. Allergenic activity is lost following

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1 proteolytic digestion (see review by BALDO, SUTTON &  
2 WRIGLEY, 1982), but is resistant to heat treatment, e.g.  
3 100°C for 30 minutes at neutral pH (MARSH et al., 1966).

5        Lol pI is defined as an allergen because of its  
6 ability to bind to specific IgE in sera of  
7 ryegrass-sensitive patients, to act as an antigen in IgG  
8 responses, and to trigger T-cell responses. The  
9 allergenic properties have been assessed by direct skin  
10 testing of grass pollen-sensitive patients. The results  
11 showed that 84% had a skin sensitivity to Lol pI  
12 (FREIDHOFF et al., 1986), demonstrating the primary  
13 importance of this protein as the major allergen.  
14 Furthermore, 95% of patients demonstrated to be grass  
15 pollen-sensitive possessed specific IgE antibody that  
16 bound to Lol pI, as demonstrated by immunoblotting (FORD &  
17 BALDO, 1986).

19        Substantial allergenic cross-reactivity between  
20 grass pollens has been demonstrated using an IgE-binding  
21 assay, the radioallergo-sorbent test (RAST), for example,  
22 as described by MARSH (1970) and LOWENSTEIN (1978).

23

24        The immunochemical relationships of Lol pI with  
25 other grass pollen antigens have been demonstrated using  
26 both polyclonal and monoclonal antibodies (e.g. SMART &  
27 KNOX, 1979; SINGH & KNOX, 1985). Antibodies have been  
28 prepared to both purified proteins and IgE-binding  
29 components. These data demonstrate that the major  
30 allergen present in pollen of closely related grasses is  
31 immunochemically similar to Lol pI (SINGH & KNOX, 1985).

33        Further background information concerning grass  
34 pollen allergens can be found in the following reviews:



- 3 -

1 MARSH (1975), HOWLETT & KNOX (1984), BALDO, SUTTON &  
WRIGLEY (1982) and FORD & BALDO (1986).

3       Recent advances in biochemistry and in recombinant  
DNA technology have made it possible to synthesize  
5 specific proteins, for example, enzymes, under controlled  
conditions independent of the organism from which they are  
7 normally isolated. These biochemical synthetic methods  
employ enzymes and subcellular components of the protein  
9 synthesizing systems of living cells, either in vitro in  
cell-free systems, or in vivo in microorganisms. In  
11 either case, the principal element is the provision of a  
deoxyribonucleic acid (DNA) of specific sequence which  
13 contains the information required to specify the desired  
amino acid sequence. Such a specific DNA sequence is  
15 termed a gene. The coding relationships whereby a  
deoxyribonucleotide sequence is used to specify the amino  
17 acid sequence of a protein is well-known and operates  
according to a fundamental set of principles (see for  
19 example, WATSON, 1976).

21       A cloned gene may be used to specify the amino acid  
sequence of proteins synthesized by in vitro systems.  
23 DNA-directed protein synthesizing systems are  
well-established in the art. Single-stranded DNA can be  
25 induced to act as messenger RNA (mRNA) in vitro, thereby  
resulting in high fidelity translation of the DNA sequence.  
27

It is now possible to isolate specific genes or  
29 portions thereof from higher organisms, such as plants,  
and to transfer the genes or DNA fragments in a suitable  
31 vector, such as lambda-gt 11 phage, to microorganisms such  
as bacteria e.g. Escherichia coli. The transferred gene  
33 is replicated and propagated as the transformed  
microorganism replicates. Consequently, the transformed

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1 microorganism is endowed with the capacity to make the  
desired protein or gene which it encodes, for example, and  
3 enzyme, and then passes on this capability to its  
progeny. See, for example, Cohen and Boyer, U.S. Patent  
5 Nos. 4,237,224 and 4,468,464. The bacterial clones  
containing the recombinant phage are screened for the  
7 particular gene product (protein) by means of specific  
antibodies.

9

In accordance with the present invention, the gene  
11 encoding Lol pI is cloned and thereby permitting the large  
scale production of recombinant allergen.

13

Accordingly, one aspect of the present invention  
15 relates to a recombinant vector comprising a DNA sequence  
encoding a protein displaying allergenic activity from  
17 pollen of a grass species. More particularly, the grass  
species belongs to the family Poaceae (Gramineae), and  
19 even more particularly, to the genus Lolium. Still even  
more particularly, the allergenic protein is characterized  
21 as being immunologically cross-reactive with antibody to  
Lol pI protein of Lolium perenne pollen, namely:

23

Pooid (festucoid) grasses. GROUP 1: Triticanae:  
25 Bromus inermis, smooth brome; Agropyron repens, English  
couch; A.cristatum; Secale cereale rye Triticum  
27 aestivum, wheat. GROUP 2: Poanae: Dactylis glomerata,  
orchard grass or cocksgoot; Festuca elatior, meadow  
29 fescue; Lolium perenne, perennial ryegrass;  
L.multiflorum, Italian ryegrass; Poa pratensis, Kentucky  
31 bluegrass; P.compressa, flattened meadow grass; Avena  
sativa, oat; Holcus lanatus, velvet grass or Yorkshire  
33 fog; Anthoxanthum odoratum, sweet vernal grass;  
Arrhenatherum elatius, oat grass; Agrostis alba, red

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1 top; Phleum pratense, timothy; Phalaris arundinacea,  
reed canary grass. Panicoid grass, Paspalum notatum,  
3 Bahia grass, Andropogonoid grasses: Sorghum halepensis,  
Johnson grass.

5

Another aspect of the present invention relates to a  
7 recombinant vector comprising a DNA sequence encoding the  
allergenic protein Lol pI of ryegrass, Lolium perenne, L.  
9 pollen, or a derivative or homologue thereof. More  
particularly, the present invention relates to a  
11 recombinant DNA molecule comprising a eukaryotic or  
prokaryotic origin of replication, a detectable marker, a  
13 DNA sequence encoding the Lol pI allergenic protein or a  
derivative or a homologue thereof or an allergenic protein  
15 cross-reactive with said Lol pI protein or its derivatives  
or homologues and optionally a promoter sequence capable  
17 of directing transcription of said allergenic protein.

19 Yet another aspect of the present invention  
contemplates a method for producing recombinant Lol pI or  
21 a derivative or homologue thereof or an allergenic protein  
immunologically reactive to antibodies to Lol pI or a  
23 derivative or homologue thereof, comprising culturing an  
organism containing a replicable recombinant DNA molecule,  
25 said molecule comprising a promoter capable of expression  
in said organism, the gene encoding Lol pI or its  
27 derivative or homologue or immunologically related protein  
of Lol pI located downstream of and transcribed from said  
29 promoter, a selectable marker and a DNA vehicle containing  
a prokaryotic or eukaryotic origin of replication, under  
31 conditions and for a time sufficient for said recombinant  
DNA molecule to be stably maintained and direct the  
33 synthesis of Lol pI or its derivative or homologue.

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1 In yet another aspect of the present invention,  
there is provided non-native (i.e., recombinant or  
3 chemically synthesized) Lol pI or its derivative or  
homologue or a non-native allergenic protein  
5 immunologically cross-reactive to antibodies to Lol pI or  
its derivative or homologue .

7

Still yet another aspect of the present invention  
9 relates to antibodies to non-native Lol pI or a derivative  
or homologue thereof.

11

In still yet another aspect of the present  
13 invention, there is provided a method for detecting an  
antibody to an allergenic protein from pollen of the  
15 family Poaceae (Gramineae) in serum or other biological  
fluid comprising contacting said serum or fluid with  
17 recombinant Lol pI or its antigenic derivative for a time  
and under conditions sufficient for an antibody - Lol pI  
19 complex to form and subjecting said complex to a detecting  
means.

21

Another aspect of the present invention relates to a  
23 recombinant DNA molecule comprising a ryegrass pollen  
promoter sequence or homologue or degenerate form thereof  
25 located on said molecule and further having one or more  
restriction sites down stream of said promoter such that a  
27 nucleotide sequence inserted into one or more of these  
sites is transcribeable in the correct reading frame.

29

In one preferred embodiment, the recombinant DNA  
31 molecule comprises the promoter directing synthesis of  
Lol pI from pollen of ryegrass, Lolium perenne L. and is  
33 thereby a developmentally regulated, pollen specific,  
expression vector.

35

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1           A further aspect of the present invention  
contemplates a method for inducing nuclear male sterility  
3 in plants of the family Poaceae comprising the steps of:  
a) developing a plant carrying a recombinant DNA  
5 molecule comprising the ryegrass pollen promoter sequence  
or homologue or degenerate form thereof located on said  
7 molecule and a nucleotide sequence encoding a polypeptide  
having a deleterious function in cells derived from the  
9 family Poaceae, said nucleotide sequence transcribeable  
from said promoter, and said recombinant DNA molecule  
11 stably contained in pollen producing cells, and,  
b) growing said plants under conditions and for a  
13 time sufficient for their developmental stage to cause  
expression of said nucleotide sequence from said promoter  
15 thereby producing the polypeptide having a deleterious  
function on said pollen producing cells such that pollen  
17 formation is inhibited or said pollen is inactive.

19           Further features of the present invention will be  
better understood from the following detailed description  
21 of the preferred embodiments of the invention in  
conjunction with the appended figures.

23

Standard biochemical nomenclature is used herein in  
25 which the nucleotide bases are designated as adenine (A);  
thymine (T); guanine (G) and cytosine (C). Other  
27 abbreviations include:-

29 BSA	Bovine serum albumin.
DEPC	Diethyl pyrocabonate.
31 DNA	Deoxyribonucleic acid.
DTT	Dithiothreitol.
33 EDTA	Disodium ethylene diamine tetra-acetate.
IPTG	Isopropyl-thio-beta-D-galactopyranoside.

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- 1  
LB MEDIUM      Luria-Bertani medium (1% (w/v) Bactotryptone,  
3                      1% (w/v) NaCl & 0.5% (w/v) Bacto-yeast  
                    extract in water to pH 7.5).
- 5  
LIGATION BUFFER (10 x solution)  
7                      0.66 M Tris Cl (pH 7.5) 50mM Mg Cl<sub>2</sub>, 50mM  
                    DTT, 10mM ATP.
- 9  
LiCl              Lithium chloride.
- 11 PEG             Polyethylene glycol.  
     pfu            Plaque forming units.
- 13 PMSF            Phenylmethylsulphonylfluoride.
- 15 SAMPLE BUFFER  
                    50mM Tris-Cl, pH 6.8, 1.5% (w/v) SDS, 50mM  
17                      DTT, 4M Urea, 1m MPMSF.
- 19 SDS             Sodium dodecyl sulfate.
- 21 SM BUFFER      Phage storage buffer (0.1M NaCl, MgSO<sub>4</sub>·7H<sub>2</sub>O,  
                    50mM Tris HCl pH 7.5, 2% (w/v) gelatin).
- 23  
SSC                20 x solution of 3M NaCl, 0.3M Na<sub>3</sub> citrate,  
25                      pH 7.0.
- SSPE            (0.15M NaCl, 10mM Sodium Phosphate pH 7.7, 1mM
- 27 EDTA            Ethylene diamine tetra-acetic acid
- TBS             Tris buffered saline (50mM Tris pH 7.5, 150  
29                      mM NaCl).
- X-gal            5-Bromo-4-chloro 3-indolyl  
31                      beta-D-galactopyranoside.

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1 In the accompanying figures:

3 Figure 1 shows the identification of Lol pI as the  
principal allergen of ryegrass pollen by SDS-PAGE and  
5 Western Blotting.

7 Lanes 1-3. SDS-PAGE analysis of total ryegrass  
pollen proteins and isolated Lol pI allergen, stained with  
9 Coomassie blue for proteins. Lane 1, total pollen  
proteins; Lane 2, isoallergen of Lol pI; Lane 3, Lol pI.  
11

Lanes 4-6. Western blot of proteins shown in Lanes  
13 1-3, showing specific binding of monoclonal antibody  
FMC-A1 to the Lol pI allergen. This antibody was used to  
15 screen the cDNA library to select the Lol pI allergen  
clones. Lane 4, molecular markers; Lane 5, isoallergen;  
17 Lane 6, Lol pI, 32 kD.

Figure 2 shows screening of cDNA library of ryegrass  
19 pollen to select the specific clones which express Lol pI  
protein in lambda-gt 11 vector. (a,b): Plaque - lifts of  
21  $10^2 - 10^3$  recombinant phages treated with specific  
antibody FMC-A1, with (a) cDNA clone 6; (b) cDNA clone  
23 12; (c,d) Re-screening of cDNA clone 12 with (c)  
monoclonal antibody FMC-a1, (d) specific IgE from ryegrass  
25 pollen-sensitive patients' sera. Recombinant phage  
containing the specific allergen DNA insert are detected  
27 by these methods. The antibodies detect all clones which  
contain the antigenic determinants of Lol pI, while IgE  
29 binds to clones containing the allergenic determinants of  
Lol pI. All clones were monoclonal antibody FMC-A1  
31 positive, as this is the basis of the screen, while a  
proportion bind to IgE, as with clone 12 here.

33

Figure 3 shows analysis of E. coli fusion protein  
35 for identity with Lol pI.

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1

Lanes 1-4: SDS-PAGE stained with Coomassie blue for  
3 proteins; Lane 1, lambda-gt 11 non-recombinant lysogen  
extract; Lane 2, lambda-l2RL8 recombinant lysogen  
5 extract; Lane 3, E. coli beta-galactosidase pure  
protein; Lane 4, molecular weight markers, the 96KD  
7 marker is indicated by a star.

Lanes 5-7: Transblots on nitrocellulose membrane;  
9 Lane 5 and 8, molecular weight markers; Lane 6,  
lambda-l2RL8 recombinant lysogen extract, a indicates a  
11 fusion protein identified by binding with FMC-A1,  
molecular weight a>c. Lane 7, lambda-6RL2 recombinant  
13 lysogen extract, B indicates fusion protein as identified  
by binding of FMC-A1, molecular weight b>c. The higher  
15 molecular weights of a and b over c indicates the  
insertion of cDNA into the gt 11 genome at the lacZ site.

17

Figure 4 shows analysis of tissue and organ  
19 specificity of Lol pI gene in ryegrass. (a) Slot blot.  
2ug each of total RNA isolated from pollen (p), leaf (l),  
21 roots (r) and hydrated seeds (s) were slot blotted onto  
nitrocellulose membrane. Hybridization with redioactive  
23 probes for clones 6 and 12 (p6, pl2) occurs with pollen,  
but there is a total absence of hybridization with the  
25 othe rtissue RNA. (b) Northern blot. Total RNA isolated  
from these ryegrass tissues were separated  
27 electrophoretically in a denaturing agarose gel, and  
transferred to nitrocellulose membrane, and probed with  
29 p6. Hybridization occurs with the pollen sample only,  
other tissue RNA showing absence of hybridization. This  
31 evidenc eshows that Lol pI gene is expressed only in  
pollen.



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1

Figure 5 shows a 1240 base pair DNA sequence  
3 representing the cDNA clone 12R

5 Figure 6 shows the reaction of recombinant allergen  
pGEX-12R (Lol pI) with IgE from pooled allergic sera. The  
7 cultures of pGEX and pGEX-12R were grown overnight and  
then diluted 1:10 in broth and grown for 2h at 37°C. They  
9 were induced with IPTG, and grown for 1h at 37°C. The  
bacteria were pelleted and resuspended in PBS to 1/20 the  
11 volume of culture media. The bacteria were lysed by  
freeze thaw and sonication. Following that an equal  
13 volume of SDS gel sample buffer was added, and samples  
boiled for 3 min, before loading them onto a 10-15%  
15 gradient SDS-PAGE. The separated proteins were  
transferred onto nitrocellulose membrane, and these blots  
17 were processed for identification of IgE-binding proteins  
using pooled sera from allergic patients. <sup>125</sup>I-labelled  
19 anti-human IgE antibodies (Kallestad Labs USA) were used  
as probe. Figure 6 shows a typical autoradiograph in  
21 which lane 1 shows a vector control in which no IgE  
binding is present, while lanes 2,3 and 4 show expression  
23 of recombinant Lol pI in bacterial cultures infected with  
pGEX-12R.  
25

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1        Figure 7 A,B and C shows antigenic and allergenic  
similarity of proteins homologous with Lol pI in a panel  
3 of 17 different grasses. Proteins were resolved by  
SDS-PAGE from mature pollen as follows: lane a: molecular  
5 weight markers; 1, Bromus inermis; 2, Agropyron  
cristatum; 3, Secale cereale; 4, Dactylis glomerata; 5,  
7 Festuca elatior; 6, Lolium perenne; 7, L. multiflorum;  
8, Poa compressa; 9, Avena sativa; 10, Holcus lanatus;  
9 11, Anthoxanthum odoratum; 12, Agrostis alba; 13, Phleum  
pratense; 14, Phalaris arundinacea; 15, Cynodon  
11 dactylon; 16, Sorghum halepensis; 17, Zea mays. 7A shows  
Coomassie blue stained proteins in SDS-PAGE gel. 7B shows  
13 western blot probed with monoclonal antibody FMC-A1  
specific for Lol pI, showing antigenic similarity of Lol  
15 pI and homologous allergens in related grasses, except for  
lane 15, Cynodon dactylon. 7C shows western blot probed  
17 with pooled allergic human sera and anti-IgE antibodies,  
confirming that Lol pI and its homologous allergens in  
19 other grasses are the immunodominant allergen of grass  
pollen.

21

Figure 8A shows a comparison of allergenic activity  
23 of native and recombinant Lol pI protein. Sera from 28  
different patients, some of whom are allergic to grass  
25 pollen, were used to compare the IgE binding of native and  
recombinant Lol pI protein. For native Lol pI, a  
27 reference standard sample was purchased from the National  
Institutes of Health (NIAID), Bethesda, USA. This sample  
29 was diluted in 1% (w/v) BSA solution, and 0.5ug was  
dot-blotted onto nitrocellulose membrane, and the blots  
31 used for IgE-binding assay. For testing IgE-binding to  
recombinant Lol pI protein, the clone lambda-gt 11 -12R  
33 was expressed in host E. coli cells. The plaque lifts  
were used in a similar way to dot blots for testing IgE

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1 binding. Both the plaque lifts and dot blots were  
incubated overnight in 1:10 dilution of allergic sera, and  
3 binding of IgE visualized using rabbit anti-human IgE  
(Dakopatts, Copenhagen, Denmark). This incubation was  
5 followed by peroxidase-conjugated goat anti-rabbit IgG,  
and then the enzyme substrate to give a colour reaction.  
7 Figure 8B is a correlation of allergenic reactivity of  
native and recombinant Lol pI.

9

Figure 9 shows restriction map of cDNA insert to  
11 lambda-gt 11 -12R, and the strategy of nucleotide  
sequencing.

13

In accordance with the present invention, there is  
15 provided the gene encoding the ryegrass pollen allergen  
Lol pI, a method for expressing same in a host cell, and  
17 more particularly organ specific (i.e., pollen), thereby  
providing a source of recombinant Lol pI and the promoter  
19 of the Lol pI gene directing developmental regulation of  
Lol pI or any genetic sequence placed downstream thereof.

21

The original source of the genetic material is fresh  
23 ryegrass pollen from Lolium perenne L., collected from  
field sources near Melbourne, Australia and bulk collected  
25 pollen from a supplier (Greer Laboratories, Lenoir, NC).  
These sources of pollen are not intended to limit the  
27 scope of the invention since they only represent one  
convenient supply of the pollen. The present invention  
29 can be practised using pollen from any location. Figure 1  
shows the identification of Lol pI as the principle  
31 allergen of ryegrass pollen.

33

"Gene", is used, in respect of the present  
invention, in its broadest sense and refers to any  
35 contiguous sequence of nucleotides, the transcription of  
which, leads to a mRNA molecule, whether or not said mRNA

- 14 -

1 molecule is translatable into a polypeptide or protein.  
The gene encoding Lol pI means the nucleotide sequence  
3 encoding the entire polypeptide or derivatives or  
homologues of said polypeptide which may contain amino  
5 acid substitutions, deletions or additions. Similarly, in  
relation to the carbohydrate portion of said polypeptide,  
7 derivatives include substitutions, deletions or additions  
to said carbohydrate moiety. The Lol pI gene also refers  
9 to a cDNA complementary to the mRNA corresponding to the  
full or partial length of the Lol pI polypeptide.

11 Accordingly, it is within the scope of the present  
invention to encompass Lol pI and its amino acid and/or  
13 carbohydrate derivatives and to nucleotide sequences,  
including DNA, cDNA and mRNA and to the homologue or  
15 degenerate forms thereof, encoding said Lol pI or said  
derivatives. It is further in accordance with the present  
17 invention to include molecules such as polypeptides fused  
to Lol pI or its derivatives or to nucleotide sequences  
19 contiguous to the Lol pI- and/or derivative-encoding  
nucleotide sequences. For example, for some aspects of  
21 the present invention, it is desirable to produce a fusion  
protein comprising Lol pI or its derivative and an amino  
23 acid sequence from another polypeptide or protein,  
examples of the latter being enzymes such as  
25 beta-galactosidase, phosphatase, urease and the like.

Most fusion proteins are formed by the expression of a  
27 recombinant gene in which two coding sequences have been  
joined together such that their reading frames are in  
29 phase. Alternatively, polypeptides can be linked in vitro  
by chemical means. All such fusion protein or hybrid  
31 genetic derivatives of Lol pI or its encoding nucleotide  
sequence are encompassed by the present invention.

33 Furthermore, by homologues and derivatives of Lol pI is  
meant to include synthetic

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1 derivatives thereof. The nucleotide sequence as  
2 elucidated herein, can be used to generate any number of  
3 peptides or polypeptides by chemical synthesis, such as  
4 solid phase synthesis, by well known methods. All such  
5 chemically synthesized peptides are encompassed by the  
6 present invention. Accordingly, the present invention  
7 extends to non-native Lol pI, and its derivatives,  
8 homologues and immunological relatives made by recombinant  
9 means or by chemical synthesis. Furthermore, the present  
10 invention extends to proteins, polypeptides or peptides  
11 corresponding in whole or part to the nucleotide coding  
12 sequence given in Figure 5 or to degenerate or homologue  
13 forms thereof.

15 It is also within the scope of the present invention  
16 to include allergenic proteins immunologically  
17 cross-reactive with antibodies to Lol pI or its  
18 derivatives or homologues. "Immunologically  
19 cross-reactive" is used in its broadest sense and refers  
20 generally to a protein capable of detectable binding to an  
21 antibody, the latter being specific to Lol pI or to  
22 derivatives or homologues of Lol pI. Such an  
23 immunologically related allergen is referred to herein as  
24 a immunological relative of Lol pI.

25

26 The cloning of the cDNA encoding Lol pI was based on  
27 the recognition of the protein expressed by Escherichia  
28 coli transformed with lambda-gt 11 phage, using both  
29 specific monoclonal antibodies and specific serum IgE from  
30 grass pollen-sensitive patients. Two such clones are  
31 designated 6R and 12R. cDNA clones were also isolated on  
32 the basis of differential antibody binding. For example,  
33 cDNA clone 6R, was isolated on the basis that it encoded a  
34 polypeptide capable of binding to monoclonal antibodies  
35 but not IgE. Polypeptides of this type apparently lack  
36 the amino acid sequence specifying allergenicity and  
37 hence, these cDNA clones must lack the DNA sequence  
38 encoding same. Monoclonal antibodies used herein are FMC  
39 A1, A5 and A7 as described by KNOX & SINGH (1985).

- 16 -

1

For cloning the Lol pI gene or derivatives thereof,  
3 mRNA was first isolated from the mature ryegrass pollen.  
This mRNA was used as a template to synthesize double  
5 stranded complementary DNAs

7

From this cDNA library of ryegrass pollen  
recombinant phage containing the Lol pI insert were  
9 detected by screening the library with (1) specific  
monoclonal antibody FMC-A1; (2) specific IgE from sera of  
11 ryegrass-sensitive patients (Fig. 2).

13

EcoRI, linkers were then attached to both sides of  
selected clones of ds cDNA and then ligated into EcoRI,  
15 lambda-gt 11 vector arms (cut and dephosphorylated as  
purchased from Promega). The recombinant lambda-gt 11 DNA  
17 containing cDNA inserts were packaged into mature phage  
and the recombinant phage allowed to infect the E. coli  
19 host.

21

The synthesis of beta-galactosidase - recombinant  
gene fusion protein was induced by adding IPTG. The  
23 Lol pI-beta-galactosidase fusion protein was then detected  
using monoclonal antibodies which specifically recognise  
25 the epitopes on Lol pI protein.

27

This fusion protein was isolated in preparative  
amounts from bacterial lysogens, fractionated by  
29 SDS-polyacrylamide gel electrophoresis, and the proteins  
transferred to nitrocellulose membranes for probing with  
31 monoclonal antibodies (Fig. 3). These antibodies  
recognised a protein which shows a molecular weight  
33 greater than the E. coli beta-galactosidase as would be  
expected of an allergen beta-galactosidase fusion protein.

- 17 -

1

The allergenic nature of the subject proteins are characterised in part, by their binding of the reagenic IgE antibodies which are present at high levels in sera of allergic patients. The IgE binding to the epitopes on allergic proteins can be tested in a chromogenic assay in which allergens immobilized on a solid support can be visualised by sequential incubation in (1) allergic patients serum; (2) enzyme-labelled anti-IgE antibodies.

Selected cDNA clones were used to probe total RNA isolated from other ryegrass plant organs to test whether Lol pI allergen is pollen-specific or not. Slot-blotting and Northern analyses were employed (Fig. 4). No hybridization was detectable for total RNA from leaf, seed or root samples. These data indicate that Lol pI is not expressed in these other organs of the ryegrass plant.

Selected cDNA clones were ligated into both M13 and Gemini vectors for sequencing. DNA restriction fragments to be sequenced were inserted into M13 mp14 (MESSING AND VIEIRA 1982). M13 cloning and dideoxy chain termination DNA sequencing were performed as described by Bio-rad Laboratories (1980) and MESSING (1983). A similar approach is used for the cloning of allergenic proteins from pollen of other members of the family Poaceae (Gramineae) which are immunologically cross-reactive with antibodies to Lol pI or its derivatives or homologues. The sequence of the 1240 base pair cDNA clone 12R is shown in Figure 5. It is in accordance with this invention to include or degenerate forms of said sequence and/or nucleotide sequences having substantial i.e., at least 60% homology thereto.

With this knowledge in hand, a variety of expression vectors can be constructed for the production of Lol pI or its derivatives. Accordingly, another aspect of the present invention contemplates a method of producing recombinant Lol pI or its derivative or homologue or its

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1 immunological relative (as hereinbefore defined)  
2 comprising culturing an organism containing a replicable  
3 recombinant DNA molecule, said molecule comprising a  
4 promoter capable of expression in said organism, the  
5 Lol pI gene or gene encoding its derivative, homologue or  
6 immunological relative thereof, located downstream of and  
7 transcribed from said promoter, a selectable marker and a  
8 DNA vehicle containing a prokaryotic or eukaryotic origin  
9 of replication, under conditions and for a time sufficient  
10 for said recombinant DNA molecule to be stably maintained  
11 and direct the synthesis of Lol pI or its derivative,  
12 homologue or immunological relative and then isolating  
13 same.

14 "Promoter" is used in its broadest sense and refers  
15 generally to nucleotide sequence which binds RNA  
16 polymerase and directs same to the correct transcriptional  
17 start site whereupon a gene or other nucleotide sequence  
18 thereof is transcribed. As used herein, a gene or  
19 nucleotide sequence is said to be relative to the promoter  
20 meaning that said promoter directs the transcription of  
21 the gene or nucleotide sequence. The promoter is also  
22 selected on the basis of its ability to function in a  
23 particular host. The following description relates to  
24 developing prokaryotic expression vectors capable of  
25 expressing the Lol pI gene or a gene encoding its  
26 derivative, homologue or immunological relative, thereof.  
27 Similar principles apply for the construction of  
28 eukaryotic vectors. In this description, reference to the  
29 Lol pI gene also includes reference to genes encoding  
30 derivative, homologues or immunological relatives of  
31 Lol pI.



- 19 -

1

In constructing suitable prokaryotic expression  
vectors, transcription termination sequences are desirable  
to prevent potential readthrough by the RNA polymerase.  
To avoid any potential interference with the transcription  
terminators, one skilled in the art can eliminate the 3'  
non-coding region of the Lol pI gene. Concurrently, one  
can substitute other known transcription terminators, for  
example, the bacteriophage lambda terminator. Thus, the  
present invention is in no way limited to the use of any  
one prokaryotic transcription terminator. Other  
transcription terminators include, for example, the lpp  
terminator and the phage SP01 terminator. All of the  
aforementioned terminators have been previously  
characterized, are well known in the art, and can be  
constructed either synthetically or from known plasmids.

17

Expression of Lol pI activity in E. coli is in no  
way limited to the use of a particular promoter, since the  
choice of a specific promoter is not critical to the  
operability of this aspect of the present invention.

Promoters which can be substituted for the previously  
exemplified  $\lambda P_L$  promoter include, but are not limited to,  
the E. coli lactose (lac), the E. coli tryptophan (trp),  
the E. coli lipoprotein (lpp), and bacteriophage lambda P  
promoters. In addition, one or more promoters can be used  
in tandem, such as, for example, the trp and lac  
promoters, or hybrid promoters, such as the tac promoter,  
can be used to drive expression of the Lol pI gene. All  
of the aforementioned promoters have been previously  
characterized, are well known in the art, and can be  
constructed either synthetically or from known plasmids.

33

- 20 -

1 The present invention is not limited to the use of  
any particular prokaryotic replicon. Many replicons, such  
3 as those from plasmids pBR322, pACYC184, the pUC plasmids,  
and the like, are known in the art and are suitable for  
5 the construction of recombinant DNA cloning and expression  
vectors designed to drive expression of the Lol pI  
7 -encoding DNA compounds of the present invention. Neither  
is the present invention limited to the actual selectable  
9 markers present on the plasmids exemplified herein. A  
wide variety of selectable markers exist, both for  
11 eukaryotic and prokaryotic host cells, that are suitable  
for use on a recombinant DNA cloning or expression vector  
13 comprising a DNA compound (or sequence) of the present  
invention.

15

Many modifications and variations of the present illustrative DNA sequences and plasmids are possible. For example, the degeneracy of the genetic code allows for the substitution of nucleotides throughout polypeptide coding regions as well as for the substitution of the TAA or TGA translation stop signals for the TAG translational stop signal. Such sequences can be deduced from the amino acid or DNA sequence of Lol pI and can be constructed by following conventional synthetic procedures. Therefore, the present invention is no way limited to the DNA sequences and plasmids specifically exemplified.

29

The practice of this invention using prokaryotic  
31 expression vectors as well as the methods disclosed in  
this invention can be applied to a wide range of host  
33 organisms, especially Gram-negative prokaryotic organisms  
such as Escherichia coli, E. coli K12, E. coli K12 RV308,

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1 E. coli K12 HB101, E. coli K12 C600, E. coli K12 SF8,  
2 E. coli K12 RR1, E. coli K12 RR1 M15, E. coli K12 MM294,  
3 E. coli SG936, and the like. Escherichia coli SG936 is  
disclosed in BUELL et al. (1985). Two of the genetic  
5 mutations introduced in this strain, the lon and htpR  
mutations are known to promote the expression of desired  
7 proteins (see, for example, GOFF and GOLDBERG (1985).  
These mutations can be transduced into other strains of  
9 E. coli by P1 transduction according to the teaching of  
MILLER (1972).

11

Alternatively, other prokaryotes can be readily  
13 employed such as Bacillus, Pseudomonas and the like.  
Minor modifications will need to be made to the expression  
15 vector depending on the host cell employed so that the  
vector replicates, the promoter functions and the  
17 selectable marker is expressed. Such modifications would  
be routine for one skilled in the art.

19

Similar considerations apply in developing  
21 eukaryotic expression vectors and many are available for  
use in mammalian cells, yeast and fungal cells and insect  
23 cells. A convenient reference guide to developing  
eukaryotic or prokaryotic expression vectors can be found  
25 in MANIATIS et al. (1982)

27 The present invention also extends to the promoter  
of ryegrass pollen proteins, and particularly, to the  
29 promoter of the Lol pI gene. This promoter  
developmentally regulates Lol pI gene expression and is  
31 organ, i.e., pollen specific. Developmental regulation as  
used herein refers to the expression of a particular  
33 trait, in this case allergenic proteins in pollen, during  
a certain stage in a plants life cycle and non-expression  
35 during another stage.

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1 Hence, the Lol pI promoter is particularly useful in  
allowing expression of Lol pI, or any other gene or  
3 nucleotide sequence relative thereto, only during the  
development of pollen. The skilled artisan will  
5 immediately recognise the importance of such a promoter in  
selectively expressing a particular trait during pollen  
7 formation.

9 Accordingly, the present invention contemplates a  
method of inhibiting pollen development or function and  
11 thereby inducing nuclear male sterility in plants of the  
family Poaceae, and in particular Lolium perenne L.,  
13 comprising the steps of:

a) developing a plant carrying a recombinant DNA  
15 molecule comprising the ryegrass pollen promoter sequence  
or homologue or degenerate form thereof located on said  
17 molecule and a nucleotide sequence encoding a polypeptide  
having a deleterious function in cells derived from the  
19 family Poaceae, said nucleotide sequence transcribeable  
from said promoter, and said recombinant DNA molecule  
21 stably contained in pollen producing cells, and,

b) growing said plants under conditions and for a  
23 time sufficient for their developmental stage to cause  
expression of said nucleotide sequence from said promoter  
25 thereby producing the polypeptide having a deleterious  
function on said pollen producing cells such that pollen  
27 formation is inhibited or said pollen is inactive.

29 Well established methods exist for introducing  
recombinant DNA molecules into plant cells such as use of  
31 Agrobacterium plasmids and electroporation amongst  
others. By "deleterious function" in respect of a  
33 polypeptide refers to a feature of said polypeptide that  
will inhibit cell growth, cause lysis of a cell, or

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1 inhibit various functions in a cell and thereby preventing  
the normal functioning of the cell. In this case, lethal  
3 gene constructs having a deleterious function are  
contemplated which inhibit or prevent pollen formation and  
5 thereby result in a male sterile plant. Such "lethal  
genes" may encode enzymes, enzyme inhibitors, and/or toxic  
7 polypeptides, amongst other molecules. Alternatively, the  
lethal gene may encode an antisense RNA capable of  
9 inhibiting translation of a particular species of mRNA,  
the translated product thereof, being vital for pollen  
11 development.

13 Male sterile plants are particularly useful in  
developing hybrid crop varieties.

15

The Lol pI promoter is isolatable from ryegrass  
17 genomic DNA by any number of procedures including use of  
promoter probes vectors, "chromosome walking" and S1  
19 nuclease mapping and sequencing as DNA upstream of the  
transcription initiation site. All these techniques are  
21 well known to the skilled artisan. For example, using the  
cDNA clone encoding Lol pI or its derivative as probe DNA  
23 for hybridization, a fragment of DNA adjacent to or  
encompassing part or all of the Lol pI gene is cloned.  
25 The nucleotide sequence of Lol pI as determined in  
accordance with the present invention, is then used, to  
27 develop nucleotide primers at the promoter-proximal end of  
the Lol pI gene. "Chromosome walking", S1 endonuclease  
29 mapping, promoter probes will readily identify the  
promoter.

31

Accordingly, the present invention contemplates a  
33 recombinant DNA molecule comprising a ryegrass pollen  
promoter sequence, and in particular the promoter for the

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1 Lol pI gene, or homologue or degenerate form thereof  
located on said molecule and further having one or more  
3 restriction endonuclease sites downstream of said promoter  
such that nucleotide sequence inserted into one or more of  
5 these sites is transcribeable in the correct reading  
frame. As used herein, the "correct reading frame" has  
7 the same meaning as "in phase". The aforementioned DNA  
molecule will preferably also have a selectable marker  
9 thereon, such as an antibiotic or other drug resistance  
gene, such as for example gene encoding resistance to  
11 ampicillin, carbenicilin, tetracycline, streptomycin and  
the like. The recombinant molecule will further comprise  
13 a means for stable inheritance in a prokaryotic and/or  
eukaryotic cell. This can be accomplished by said  
15 recombinant molecule carrying a eukaryotic and/or a  
prokaryotic origin of replication as hereinbefore  
17 described in relation to expression vectors.

Alternatively, the recombinant molecule will carry a means  
19 for integration into a host cell genome thereby permitting  
replication of said recombinant molecule in synchrony with  
21 the replication of said host cell genome. Examples of  
preferred prokaryotic hosts include E. coli, Bacillus and  
23 Pseudomonas amongst others. Preferred eukaryotic hosts  
include cells from yeast and fungi, insects, mammals and  
25 plants. Even more preferred host cells are plants of the  
family Poaceae, and in particular of the genus Lolium,  
27 such as Lolium perenne. Accordingly in a preferred  
embodiment, the Lol pI gene promoter with a gene encoding  
29 a deleterious function positioned relative thereto will be  
carried by a recombinant DNA molecule capable of  
31 integration into the genome of cells of plants from the  
family Poaceae, or more particularly, of the genus Lolium,  
33 such as Lolium perenne. Such a recombinant DNA molecule  
is transferred to the aforementioned cells by, for

- 25 -

1 example, electroporation. Ideally, said cells are  
callus-derived cells. Said callus-derived cells  
3 transformed with said recombinant DNA molecule are then  
permitted to regenerate into whole plants. Whole plants  
5 entering the pollen development stage of its life cycle,  
permit functioning of the Lol pI gene promoter and hence,  
7 expression of the gene encoding a deleterious function.  
Consequently pollen development is inhibited or prevented  
9 and a nuclear male sterile plant results therefrom.

11 Alternatively, the Lol pI promoter will direct  
expression of a gene having advantageous functions, such  
13 as a cytokinin. All such recombinant DNA molecules are  
encompassed by the present invention.

15

The monoclonal antibodies used in the present work  
17 to screen the cDNA library for Lol pI clones showed  
cross-reactivity with allergenic proteins from pollen of  
19 various related grass species. This shows there is a  
homology between allergenic proteins produced by these  
21 pollens with Lol pI allergen supporting the applicability  
of the present invention to all related grasses. For  
23 example, this homology can be exploited to isolate DNA  
encoding other allergenic proteins without the need for  
25 protein microsequencing and oligo-nucleotide primers.  
The present invention also relates to antibodies to  
27 recombinant Lol pI and its derivatives, homologues and  
immunological relatives including its chemical synthetic  
29 derivatives In the following discussion, reference to Lol  
pI includes its derivatives, homologues and immunological  
31 relatives and chemical synthetic derivatives thereof.  
Such antibodies are contemplated to be useful in  
33 developing detection assays (immunoassays) for said Lol pI  
especially during the monitoring of a therapeutic or  
35 diagnostic regimen and in the purification of Lol pI. The  
antibodies may be

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1 monoclonal or polyclonal. Additionally, it is within the  
scope of this invention to include any second antibodies  
3 (monoclonal or polyclonal) directed to the first  
antibodies discussed above. The present invention further  
5 contemplates use of these first or second antibodies in  
detection assays and, for example, in monitoring the  
7 effect of a diagnostic or an administered pharmaceutical  
preparation. Furthermore, it is within the scope of the  
9 present invention to include antibodies to the  
glycosylated regions of Lol pI (where present), and to any  
11 molecules complexed with said Lol pI. Accordingly, an  
antibody to Lol pI encompasses antibodies to Lol pI, or  
13 antigenic parts thereof, and to any associated molecules  
(e.g., glycosylated regions, lipid regions, carrier  
15 molecules, fused proteins, and the like).

17       The Lol pI, or parts thereof, considered herein are  
purified then utilized in antibody production. Both  
19 polyclonal and monoclonal antibodies are obtainable by  
immunization with Lol pI, and either type is utilizable  
21 for immunoassays. The methods of obtaining both types of  
sera are well known in the art. Polyclonal sera are less  
23 preferred but are relatively easily prepared by injection  
of a suitable laboratory animal with an effective amount  
25 of the purified Lol pI, or antigenic parts thereof,  
collecting serum from the animal, and isolating specific  
27 sera by any of the known immunoabsorbent techniques.  
Although antibodies produced by this method are utilizable  
29 in virtually any type of immunoassay, they are generally  
less favored because of the potential heterogeneity of the  
31 produce.

33       The use of monoclonal antibodies in an immunoassay  
is particularly preferred because of the ability to



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1 produce them in large quantities and the homogeneity of  
the product. The preparation of hybridoma cell lines for  
3 monoclonal antibody production derived by fusing an  
immortal cell line and lymphocytes sensitized against the  
5 immunogenic preparation can be done by techniques which  
are well known to those who are skilled in the art. (See,  
7 for example, DOUILLARD, and HOFFMAN (1981) and KOHLER and  
MILSTEIN (1975; 1976).

9

Unlike preparation of polyclonal sera, the choice of  
11 animal is dependent on the availability of appropriate  
immortal lines capable of fusing with lymphocytes. Mouse  
13 and rat have been the animals of choice in hybridoma  
technology and are preferably used. Humans can also be  
15 utilized as sources for sensitized lymphocytes if  
appropriate immortalized human (or nonhuman) cell lines  
17 are available. For the purpose of the present invention,  
the animal of choice may be injected with from about  
19 0.1 mg to about 20 mg of the purified Lol pI, or parts  
thereof. Usually the injecting material is emulsified in  
21 Freund's complete adjuvant. Boosting injections may also  
be required. The detection of antibody production can be  
23 carried out by testing the antisera with appropriately  
labelled antigen. Lymphocytes can be obtained by removing  
25 the spleen or lymph nodes of sensitized animals in a  
sterile fashion and carrying out fusion. Alternatively,  
27 lymphocytes can be stimulated or immunized in vitro, as  
described, for example, in READING (1982).

29

A number of cell lines suitable for fusion have been  
31 developed, and the choice of any particular line for  
hybridization protocols is directed by any one of a number  
33 of criteria such as speed, uniformity of growth  
characteristics, deficiency of its metabolism for a  
35 component of the growth medium, and potential for good  
fusion frequency.

- 28 -

1

Intraspecies hybrids, particularly between like strains, work better than interspecies fusions. Several cell lines are available, including mutants selected for the loss of ability to secrete myeloma immunoglobulin.

7

Cell fusion can be induced either by virus, such as Epstein-Barr or Sendai virus, or polyethylene glycol.

9 Polyethylene glycol (PEG) is the most efficacious agent for the fusion of mammalian somatic cells. PEG itself may be toxic for cells, and various concentrations should be tested for effects on viability before attempting fusion. The molecular weight range of PEG may be varied from 1000 to 6000. It gives best results when diluted to from about 20% to about 70% (w/w) in saline or serum-free medium.

Exposure to PEG at 37°C for about 30 seconds is preferred in the present case, utilizing murine cells. Extremes of temperature (i.e., about 45°C) are avoided, and preincubation of each component of the fusion system at 37°C prior to fusion can be useful. The ratio between lymphocytes and malignant cells is optimized to avoid cell fusion among spleen cells and a range of from about 1:1 to about 1:10 is commonly used.

25

The successfully fused cells can be separated from the myeloma line by any technique known by the art. The most common and preferred method is to chose a malignant line which is hyposanthine Guanine Phosphoribosyl Transferase (HGPRT) deficient, which will not grow in an aminopterin-containing medium used to allow only growth of hybrids and aminopterin-containing medium used to allow only growth of hybrids and which is generally composed of hyposanthine  $1.10^{-4}M$ , aminopterin  $1 \times 10^{-5}M$ , and thymidine  $3 \times 10^{-5}M$ , commonly known as the HAT medium. The fusion

- 29 -

1 mixture can be grown in the HAT-containing culture medium  
immediately after the fusion 24 hours later. The feeding  
3 schedules usually entail maintenance in HAT medium for two  
weeks and then feeding with either regular culture medium  
5 or hyposanthine, thymidine-containing medium.

7       The growing colonies are then tested for the  
presence of antibodies that recognize the antigenic  
9 preparation. Detection of hybridoma antibodies can be  
performed using an assay where the antigen is bound to a  
11 solid support and allowed to react to hybridoma  
supernatants containing putative antibodies. The presence  
13 of antibodies may be detected by "sandwich" techniques  
using a variety of indicators. Most of the common methods  
15 are sufficiently sensitive for use in the range of  
antibody concentrations secreted during hybrid growth.  
17

Cloning of hybrids can be carried out after 21-23  
19 days of cell growth in selected medium. Cloning can be  
performed by cell limiting dilution in fluid phase or by  
21 directly selecting single cells growing in semi-solid  
agarose. For limiting dilution, cell suspensions are  
23 diluted serially to yield a statistical probability of  
having only one cell per well. For the agarose technique,  
25 hybrids are seeded in a semisolid upper layer, over a  
lower layer containing feeder cells. The colonies from  
27 the upper layer may be picked up and eventually  
transferred to wells.

29

Antibody-secreting hybrids can be grown in various  
31 tissue culture flasks, yielding supernatants with variable  
concentrations of antibodies. In order to obtain higher  
33 concentrations, hybrids may be transferred into animals to  
obtain inflammatory ascites. Antibody-containing ascites

- 30 -

1 can be harvested 8-12 days after intraperitoneal  
injection. The ascites contain a higher concentration of  
3 antibodies but include both monoclonals and  
immunoglobulins from the inflammatory ascites. Antibody  
5 purification may then be achieved by, for example,  
affinity chromatography.

7

The presence of Lol pI contemplated herein, or  
9 antibodies specific for same, in a patient's serum, plant  
or mammalian tissue or tissue extract, can be detected  
11 utilizing antibodies prepared as above, either monoclonal  
or polyclonal, in virtually any type of immunoassay. A  
13 wide range of immunoassay techniques are available as can  
be seen by reference to U.S. Patent No. 4,016,043,  
15 4,424,279 and 4,018,653. This, of course, includes both  
single-site and two-site, or "sandwich", assays of the  
17 non-competitive types, as well as in the traditional  
competitive binding assays. Sandwich assays are among the  
19 most useful and commonly used assays and are favoured for  
use in the present invention. A number of variations of  
21 the sandwich assay technique exist, and all are intended  
to be encompassed by the present invention. Briefly, in a  
23 typical forward assay, an unlabelled antibody is  
immobilized in a solid substrate and the sample to be  
25 tested brought into contact with the bound molecule. After  
a suitable period of incubation, for a period of time  
27 sufficient to allow formation of an antibody-antigen  
secondary complex, a second antibody, labelled with a  
29 reporter molecule capable of producing a detectable signal  
is then added and incubated, allowing time sufficient for  
31 the formation of a tertiary complex of  
antibody-antigen-labelled antibody (e.g. antibody Lol pI  
33 antibody). Any unreacted material is washed away, and the  
presence of the antigen is determined by observation of a

- 31 -

1 signal produced by the reporter molecule. The results may  
either be qualitative, by simple observation of the  
3 visible signal, or may be quantitated by comparing with a  
control sample containing known amounts of hapten.  
5 Variations on the forward assay include a simultaneous  
assay, in which both sample and labelled antibody are  
7 added simultaneously to the bound antibody, or a reverse  
assay in which the labelled antibody and sample to be  
9 tested are first combined, incubated and then added  
simultaneously to the bound antibody. These techniques  
11 are well known to those skilled in the art, including any  
minor variations as will be readily apparent.  
13

Although the following discussion is concerned with  
15 detecting Lol pI, it is equally applicable to detecting  
antibodies to Lol pI and it is intended to be sufficient  
17 description thereof. In the typical forward sandwich  
assay, a first antibody having specificity for Lol pI, or  
19 antigenic parts thereof, contemplated in this invention,  
is either covalently or passively bound to a solid  
21 surface. The solid surface is typically glass or a  
polymer, the most commonly used polymers being cellulose,  
23 polyacrylamide, nylon, polystyrene, polyvinyl chloride or  
polypropylene. The solid supports may be in the form of  
25 tubes, beads, discs of microplates, or any other surface  
suitable for conducting an immunoassay. The binding  
27 processes are well-known in the art and generally consist  
of cross-linking covalently binding or physically  
29 adsorbing, the polymer-antibody complex is washed in  
preparation for the test sample. An aliquot of the sample  
31 to be tested is then added to the solid phase complex and  
incubated at 25°C for a period of time sufficient to allow  
33 binding of any subunit present in the antibody. The

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1 incubation period will vary but will generally be in the  
2 range of about 2-40 minutes. Following the incubation  
3 period, the antibody subunit solid phase is washed and  
4 dried and incubated with a second antibody specific for a  
5 portion of the hapten. The second antibody is linked to a  
6 reporter molecule which is used to indicate the binding of  
7 the second antibody to the hapten.

8 By "reporter molecule," as used in the present  
9 specification, is meant a molecule which, by its chemical  
10 nature, provides an analytically identifiable signal which  
11 allows the detection of antigen-bound antibody. Detection  
12 may be either qualitative or quantitative. The most  
13 commonly used reporter molecules in this type of assay are  
14 either enzymes, fluorophores or radionuclide containing  
15 molecules (i.e. radioisotopes). In the case of an enzyme  
16 immunoassay, an enzyme is conjugated to the second  
17 antibody, generally by means of glutaraldehyde or  
18 periodate. As will be readily recognized, however, a wide  
19 variety of different conjugation techniques exist, which  
20 are readily available to the skilled artisan. Commonly  
21 used enzymes include horseradish peroxidase, glucose  
22 oxidase, beta-galactosidase and alkaline phosphatase,  
23 amongst others. The substrates to be used with the  
24 specific enzymes are generally chosen for the production,  
25 upon hydrolysis by the corresponding enzyme, of a  
26 detectable color change. For example, p-nitrophenyl  
27 phosphate is suitable for use with alkaline phosphatase  
28 conjugates; for peroxidase conjugates,  
29 1,2-phenylenediamine, 5-aminosalicylic acid, or toluidine  
30 are commonly used. It is also possible to employ  
31 fluorogenic substrates, which yield a fluorescent product  
32 rather than the chromogenic substrates noted above. In  
33 all cases, the enzyme-labelled antibody is added to the

- 33 -

1 first antibody hapten complex, allowed to bind, and then  
the excess reagent is washed away. A solution containing  
3 the appropriate substrate is then added to the tertiary  
complex of antibody-antigen-antibody. The substrate will  
5 react with the enzyme linked to the second antibody,  
giving a qualitative visual signal, which may be further  
7 quantitated, usually spectrophotometrically, to give an  
indication of the amount of hapten which was present in  
9 the sample. "Reporter molecule" also extends to use of  
cell agglutination or inhibition of agglutination such as  
11 red blood cells on latex beads, and the like.

13 Alternately, fluorescent compounds, such as  
florescein and rhodamine, may be chemically coupled to  
15 antibodies without altering their binding capacity. When  
activated by illumination with light of a particular  
17 wavelength, the fluorochrome-labelled antibody adsorbs the  
light energy, inducing a state of excitability in the  
19 molecule, followed by emission of the light at a  
characteristic color visually detectable with a light  
21 microscope. As in the EIA, the fluorescent labelled  
antibody is allowed to bind to the first antibody-hapten  
23 complex. After washing off the unbound reagent, the  
remaining tertiary complex is then exposed to the light of  
25 the appropriate wavelength, the fluorescence observed  
indicates the presence of the hapten of interest.  
27 Immunofluorescence and EIA techniques are both very well  
established in the art and are particularly preferred for  
29 the present method. However, other reporter molecules,  
such as radioisotope, chemiluminescent or bioluminescent  
31 molecules, may also be employed. It will be readily  
apparent to the skilled technician how to vary the  
33 procedure to suit the required purpose. It will also be  
apparent that the foregoing can be used to detect directly  
35 or indirectly (i.e., via antibodies) the Lol pI of this  
invention.

- 34 -

1

Accordingly, one aspect of the present invention  
contemplates a method of detecting Lol pI or a derivative  
or homologue thereof or a allergenic protein  
immunologically reactive with said Lol pI or its  
derivative or homologue in serum, tissue extract, plant  
extract or other biologically fluid comprising the steps  
of contacting said serum, extract or fluid to be tested  
with an antibody to Lol pI for a time and under conditions  
sufficient for an allergenic protein-antibody complex to  
form and subjecting said complex to a detecting means.

The present invention also contemplates a method of  
detecting an antibody to an allergenic protein from pollen  
of the family Poaceae (Gramineae) in serum or other  
biological fluid comprising contacting said serum or fluid  
with recombinant Lol pI or its antigenic derivative for a  
time and under conditions sufficient for an antibody - Lol  
pI complex to form and subjecting said complex to a  
detecting means. The latter complex may be detected by  
the Lol pI having attached thereto a reporter molecule or  
by addition of a second antibody labelled with a reporter  
molecule.

23

Accordingly, the present invention is also directed  
to a kit for the rapid and convenient assay for antibodies  
to Lol pI or its derivatives, homologues or immunological  
relatives in mammalian body fluids (e.g. serum, tissue  
extracts, tissue fluids), in vitro cell culture  
supernatants, and cell lysates. The kit is  
compartmentalized to receive a first container adapted to  
contain recombinant Lol pI, or to an antigenic component  
thereof, and a second container adapted to contain an  
antibody to Lol pI said antibody being labelled with a  
reporter molecule capable of giving a detectable signal as



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1 hereinbefore described. If the reporter molecule is an  
enzyme, then a third container adapted to contain a  
3 substrate for said enzyme is provided. In an exemplified  
use of the subject kit, a sample to be tested is contacted  
5 tot he contents of the first container for a time and  
under conditions for an antibody, if present, to bind to  
7 Lol pI in said first container. If Lol pI of the first  
container has bound to antibodies in the test fluid, the  
9 antibodies of the second container will bind to the  
secondary complex to form a tertiary complex and, since  
11 these antibodies are labelled with a reporter molecule,  
when subjected to a detecting means, the tertiary complex  
13 is detected. Therefore, one aspect of the present  
invention is a kit for the detection of antibodies to a  
15 protein having allergenic properties, said protein from  
pollen of the family Poaceae (Gramineae), the kit being  
17 compartmentalized to receive a first container adapted to  
contain recombinant Lol pI or its antigenic derivative or  
19 homologue, and a second container adapted to contain and  
antibody to Lol pI or its derivative or homologue, said  
21 antibody labelled with a reporter molecule capable of  
giving a detectable signal. The "reporter molecule" may  
23 also involve agglutination of red blood cells (RBC) on  
latex beads. In this kit the reporter molecule is a  
25 radioisotope, an enzyme, a fluorescent molecule, a  
chemiluminescent molecule, bioluminescent molecule or  
27 RBC. The kit alternatively comprises a container adapted  
to contain recombinant Lol pI or is antigenic derivative  
29 or homologue labelled with a reporter molecule capable of  
giving a detectable signal.

31

Because of the presence of allergens in the  
33 environment, hayfever and seasonal asthma continue to have  
significant morbidity and socio-economic impact on Western

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1 communities, despite advances made in their pharmacology  
and immunology. While the available spectrum of drugs,  
3 including anti-histamines and steroids have resulted in  
spectacular improvement in the treatment of allergic  
5 disease, yet they have unfortunate side-effects associated  
with longterm usage. Because of these problems, renewed  
7 interest has been shown in the immunotherapy of allergic  
disease. Immunotherapy involves the injection of potent  
9 allergen extracts to desensitize patients against allergic  
reactions (BOUSQUET, & MICHEL, 1989.) Unfortunately, the  
11 pollen preparations used as allergens are polyvalent and  
of poor quality. Consequently, concentrations used are  
13 frequently high in order to induce IgG responses, but may  
be lethal through triggering of systemic reactions,  
15 including anaphylaxis. The cloned gene product or  
synthetic peptides based on the sequence of allergens  
17 provides a safer medium for therapy since it can be  
quality controlled, characterized and standardized.

19

The precise mechanism for symptomatic relief remains  
21 hypothetical. It is established that desensitization  
therapy induces the formation of allergen-specific  
23 non-mast cell-binding IgG which blocks the combination of  
mast cell-bound IgE and allergen. This prevents mediator  
25 release, and triggering of the allergic response. Recent  
studies of ragweed pollen sensitivity showed that there is  
27 a correlation between allergen-specific IgG levels and  
relief from allergic symptoms (Lichtenstein et al.,  
29 1983). Application of reagents which can trigger  
allergen-specific IgG production during immunotherapy  
31 could significantly enhance the success rate of this  
treatment.

33

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1       Currently immunotherapy is one of the most  
frequently administered treatments in allergology, and in  
3 USA it is considered the first choice. Advantages of this  
treatment for pollen rhinitis is that treatment takes up  
5 to 3 years, while pharmacotherapy must be carried out  
during the patient's entire life time. Patients given  
7 pollen extract for immunotherapy showed a clinical benefit  
that lasted for four years after the end of treatment  
9 (GRAMMER et al., 1984.

11       Accordingly, Lol pI, its derivatives, homologues or  
immunological relatives is useful in developing a vaccine  
13 to desensitized humans to allergies due to grass pollen.

15       Accordingly, the present invention contemplates a  
method for desensitizing a human allergic to grass pollen  
17 which comprises administering to said human a  
desensitizing-effective amount of Lol pI or a derivative,  
19 homologue, or immunological relative thereof whether made  
by recombinant or synthetic means for a time and under  
21 conditions sufficient to effect desensitization of said  
human to said grass pollen.

23       The present invention, therefore, contemplates a  
25 pharmaceutical composition comprising a desensitizing  
effective amount of Lol pI or its derivatives, homologues  
27 or immunological relatives and a pharmaceutically  
acceptable carrier. The active ingredients of a  
29 pharmaceutical composition comprising Lol pI or the like  
are contemplated to exhibit excellent therapeutic  
31 activity, for example, in the desensitization of humans  
allergic to grass pollen when administered in amount

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1 which depends on the particular case. For example, from  
2 about 0.5 ug to about 20 mg per kilogram of body weight  
3 per day may be administered. Dosage regima may be  
4 adjusted to provide the optimum therapeutic response. For  
5 example, several divided doses may be administered daily  
6 or the dose may be proportionally reduced as indicated by  
7 the exigencies of the therapeutic situation. The active  
8 compound may be administered in a convenient manner such  
9 as by the oral, intraveneous (where water soluble),  
10 intramuscular, subcutaneous, intranasal, intradermal or  
11 suppository routes or implanting (eg using slow release  
12 molecules). Depending on the route of administration, the  
13 active ingredients which comprise Lol pI or the like may  
14 be required to be coated in a material to protect said  
15 ingredients from the action of enzymes, acids and other  
16 natural conditions which may inactivate said ingredients.  
17 For example, the low lipophilicity of Lol pI or the like  
18 will allow it to be destroyed in the gastrointestinal  
19 tract by enzymes capable of cleaving peptide bonds and in  
20 the stomach by acid hydrolysis. In order to administer  
21 Lol pI or the like by other than parenteral  
22 administration, they will be coated by, or administered  
23 with, a material to prevent its inactivation. For  
24 example, Lol pI or the like may be administered in an  
25 adjuvant, co-administered with enzyme inhibitors or in  
26 liposomes. Adjuvant is used in its broadest  
27 sense and includes any immune stimulating  
28 compound such as interferon. Adjuvants  
29 contemplated herein include resorcinols, non-ionic  
30 surfactants such as polyoxyethylene oleyl ether and  
31 n-hexadecyl polyethylene ether. Enzyme inhibitors include  
32 pancreatic trypsin inhibitor, diisopropylfluorophosphate  
33 (DEP) and trasylol. Liposomes include  
34 water-in-oil-in-water CGF emulsions as well as  
35 conventional liposomes.

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1       The active compounds may also be administered  
2       parenterally or intraperitoneally. Dispersions can also  
3       be prepared in glycerol, liquid polyethylene glycols, and  
4       mixtures thereof and in oils. Under ordinary conditions  
5       of storage and use, these preparations contain a  
6       preservative to prevent the growth of microorganisms.

7  
8       The pharmaceutical forms suitable for injectable use  
9       include sterile aqueous solutions (where water soluble) or  
10       dispersions and sterile powders for the extemporaneous  
11       preparation of sterile injectable solutions or  
12       dispersion. In all cases the form must be sterile and  
13       must be fluid to the extent that easy syringability  
14       exists. It must be stable under the conditions of  
15       manufacture and storage and must be preserved against the  
16       contaminating action of microorganisms such as bacteria  
17       and fungi. The carrier can be a solvent or dispersion  
18       medium containing, for example, water, ethanol, polyol  
19       (for example, glycerol, propylene glycol, and liquid  
20       polyethylene glycol, and the like), suitable mixtures  
21       thereof, and vegetable oils. The proper fluidity can be  
22       maintained, for example, by the use of a coating such as  
23       lecithin, by the maintenance of the required particle size  
24       in the case of dispersion and by the use of  
25       surfactants. The preventions of the action of  
26       microorganisms can be brought about by various  
27       antibacterial and antifungal agents, for example,  
28       parabens, chlorobutanol, phenol, sorbic acid, thimerosal,  
29       and the like. In many cases, it will be preferable to  
30       include isotonic agents, for example, sugars or sodium  
31       chloride. Prolonged absorption of the injectable  
32       compositions can be brought about by the use in the  
33       compositions of agents delaying absorption, for example,  
34       aluminum monostearate and gelatin.

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1

3 Sterile injectable solutions are prepared by  
5 incorporating the active compounds in the required amount  
7 in the appropriate solvent with various of the other  
9 ingredients enumerated above, as required, followed by  
11 filtered sterilization. Generally, dispersions are  
13 prepared by incorporating the various sterilized active  
15 ingredient into a sterile vehicle which contains the basic  
dispersion medium and the required other ingredients from  
those enumerated above. In the case of sterile powders  
for the preparation of sterile injectable solutions, the  
preferred methods of preparation are vacuum drying and the  
freeze-drying technique which yield a powder of the active  
ingredient plus any additional desired ingredient from  
previously sterile-filtered solution thereof.

17 When Lol pI or the like is suitably protected as  
described above, the active, compound may be orally  
19 administered, for example, with an inert diluent or with  
an assimilable edible carrier, or it may be enclosed in  
21 hard or soft shell gelatin capsule, or it may be  
compressed into tablets, or it may be incorporated  
23 directly with the food of the diet. For oral therapeutic  
administration, the active compound may be incorporated  
25 with excipients and used in the form of ingestible  
tablets, buccal tablets, troches, capsules, elixirs,  
27 suspensions, syrups, wafers, and the like. Such  
compositions and preparations should contain at least 1%  
29 by weight of active compound. The percentage of the  
compositions and preparations may, of course, be varied  
31 and may conveniently be between about 5 to about 80% of  
the weight of the unit. The amount of active compound in  
33 such therapeutically useful compositions in such that a  
suitable dosage will be obtained. Preferred compositions  
35 or preparations

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1 according to the present invention are prepared so that an  
2 oral dosage unit form contains between about 10 ug and  
3 2000 mg of active compound.

5       The tablets, troches, pills, capsules and the like  
6 may also contain the following: A binder such as gum  
7 gragacanth, acacia, corn starch or gelatin; excipients  
8 such as dicalcium phosphate; a disintegrating agent such  
9 as corn starch, potato starch, alginic acid and the like;  
10 a lubricant such as magnesium stearate; and a sweetening  
11 agent such a sucrose, lactose or saccharin may be added or  
12 a flavoring agent such as peppermint, oil of wintergree,  
13 or cherry flavouring. When the dosage unit form is a  
14 capsule, it may contain, in addition to materials of the  
15 above type, a liquid carrier. Various other materials may  
16 be present as coatings or to otherwise modify the physical  
17 form of the dosage unit. For instance, tablets, pills, or  
18 capsules may be coated with shellac, sugar or both. A  
19 syrup or elixir may contain the active compound, sucrose  
20 as a sweetening agent, methyl and propylparabens as  
21 preservatives, a dye and flavoring such as cherry or  
22 orange flavor. Of course, any material used in preparing  
23 any dosage unit form should be pharmaceutically pure and  
24 substantially non-toxic in the amounts employed. In  
25 addition, the active compound may be incorporated into  
26 sustained-release preparations and formulations.

27

28       As used herein "pharmaceutically acceptable carrier"  
29 includes any and all solvents, dispersion media, coatings,  
30 antibacterial and antifungal agents, isotonic and  
31 absorption delaying agents, and the like. The use of such  
32 media and agents for pharmaceutical active substances is  
33 well known in the art. Except insofar as any conventional  
media or agent is incompatible with the active ingredient,

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1 use thereof in the therapeutic compositions is  
2 contemplated. Supplementary active ingredients can also  
3 be incorporated into the compositions.

5       It is especially advantageous to formulate  
6 parenteral compositions in dosage unit form for ease of  
7 administration and uniformity of dosage. Dosage unit form  
8 as used herein refers to physically discrete units suited  
9 as unitary dosages for the mammalian subjects to be  
10 treated; each unit containing a predetermined quantity of  
11 active material calculated to produce the desired  
12 therapeutic effect in association with the required  
13 pharmaceutical carrier. The specification for the novel  
14 dosage unit forms of the invention are dictated by and  
15 directly dependent on (a) the unique characteristics of  
16 the active material and the particular therapeutic effect  
17 to be achieved, and (b) the limitations inherent in the  
18 art of compounding such an active material for the  
19 treatment of disease in living subjects having a diseased  
20 condition in which bodily health is impaired as herein  
21 disclosed in detail.

23       The principal active ingredient is compounded for  
24 convenient and effective administration in effective  
25 amounts with a suitable pharmaceutically acceptable  
26 carrier in dosage unit form as hereinbefore disclosed. A  
27 unit dosage form can, for example, contain the principal  
28 active compound in amounts ranging from 0.5  $\mu$ g to about  
29 2000 mg. Expressed in proportions, the active compound is  
30 generally present in from about 0.5  $\mu$ g to about 2000 mg/ml  
31 of carrier. In the case of compositions containing  
32 supplementary active ingredients, the dosages are  
33 determined by reference to the usual dose and manner of  
administration of the said ingredients.



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1

The present invention is further illustrated by the  
3 following non-limiting examples.

5 EXAMPLE 1

## Extraction of RNA

7

4g of fresh ryegrass pollen (collected from field  
9 sources near Melbourne and stored under liquid N<sub>2</sub>) was  
suspended in 10 ml of extraction buffer (50mM Tris buffer,  
11 pH9, 0.2M NaCl, 10mM Mg acetate), containing vanadyl  
ribonucleoside complexes to 10mM (BERGER AND BIRKENMEIER,  
13 1979) and DEPC to 0.1%. The pollen sluchs was ground in a  
mortar and pestle under liquid N<sub>2</sub> for 10-20 min to provide  
15 a homogenate in which all pollen grains are broken. The  
slurry was transferred to Nalgene centrifuge tubes with 1%  
17 (w/v) SDS, 10mM EDTA and 0.5% (w/v) N-lauroyl sarcosine.  
An equal volume of warm, high grade buffered phenol (from  
19 IBI), treated with 0.1% (w/v) hydroxyquinoline (MANIATIS  
et al., 1982) was added, and the mixture shaken for 10  
21 min. An equivalent volume of 24:1 parts  
chloroform:isoamylalcohol was added and shaking  
23 continued. Tubes were centrifuged at 15,000 rpm for 20  
min at 10°C to separate the phases and remove the  
25 insoluble material and cell debris. The aqueous phase was  
reextracted with P:C:I four times until the phenol phase  
27 remained clear, (with phase separation at 2,500 rpm for 15  
min at room temperature), and the aqueous phase was  
29 transferred to Corex centrifuge tubes. 2.5 volumes of  
100% (v/v) ethanol were added and the solution mixed by  
31 pipette, and allowed to precipitate overnight at -20°C,  
and spun at 15,000 rpm for 20 min at 0°C. The pellet was  
33 resuspended in 10 ml of EDTA to remove the  
vanadyl-ribonucleoside complexes, and LiCl added to give

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1 a final concentration of 2M. The solution was kept at 0°C  
overnight and centrifuged at 15,000 rpm for 30 min at  
3 4°C. The pellet was washed with cold 2M LiCl and 5mM EDTA  
(pH 7.3), the liquid poured off, and the pellet  
5 resuspended in 1 ml of water. The solution is heated to  
65°C, and 0.1 ml of 3M Na- acetate and 2.2 ml of ethanol  
7 were added for overnight precipitation of total RNA at  
-20°C. The pellet was washed gently with 70% (v/v)  
9 ethanol vacuum-dried, and resuspended in 0.5 ml water.  
The suspension was stored at -70°C until required for poly  
11 (A+) RNA selection.

13 One gram of ryegrass pollen contained 1 mg total  
RNA. Poly (A+) mRNA was selected by affinity  
15 chromatography on Poly (U)-Sepharose (Pharmacia)  
according to standard methods. The integrity of poly (A+)  
17 mRNA was examined in terms of its ability to act as a  
template for synthesis of single-stranded cDNA as well as  
19 its translational activity in the rabbit reticulocyte  
system.

21

## EXAMPLE 2

23 Preparation of cDNA clones

25 Synthesis of first strand cDNA was from 5ug (poly  
A+) RNA in 50 ul reaction buffer (50mM Tris buffer, pH  
27 8.3, containing 40mM KCl, 10mM MgCl<sub>2</sub>, 5mM DTT, 1mM each of  
dATP, dGTP, dTTP, and dCTP, 50 units of human placental  
29 ribonuclease inhibitor (HPRI), 5 ug of  
oligodeoxythymidylic acid primer, 80 uCi of [Alpha-<sup>32</sup>P]  
31 dCTP (3000Ci/mmol; Amersham) and 100 units of reverse  
transcriptase. The mixture was incubated for 60 min at  
33 42°C. Second strand cDNA was synthesized using the  
mRNA/cDNA hybrids as substrate, 4 units of E. coli DNA

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1 ribonuclease H to produce nicks in the mRNA template, and  
2 115 units of E. coli DNA polymerase I to catalyse the  
3 replacement of the mRNA strand by DNA. The reaction  
4 mixture was incubated sequentially at 12°C for 60 min  
5 each, and the reaction stopped by heating at 70°C for 10  
6 min. 10 units of T4 DNA polymerase were added to remove  
7 small 3' overhangs from the first strand cDNA (GUBLER AND  
8 HOFFMAN, 1983). The reaction was stopped by adding one  
9 tenth volume of 20mM EDTA and 1% (w/v) SDS. The double  
10 strand (ds) cDNA was purified by phenol/chloroform  
11 extraction followed by precipitation with ethanol.

13 In order to construct a lambda-gt 11 cDNA expression  
14 library, 500 ng of double stranded cDNA was incubated with  
15 20 units of EcoRI methylase at 37°C for 60 min.

17 1 µg of phosphorylated EcoRI linkers (5'-d[pGGAATTCC])  
18 was ligated to the double stranded cDNA in ligation buffer  
19 with 5 units of T4 DNA ligase at 15°C overnight. The  
20 EcoRI-linkered cDNA was digested with 100 units of EcoRI  
21 linkers through a Sephadryl column.

23 A 50ng of linkered cDNA was ligated to 1 µg of  
24 dephosphorylated EcoRI-cut lambda-gt 11 DNA (Promega) with  
25 2.5 units of T4 DNA ligase for 20 h at 15°C in a total  
26 volume of 10 µl. The ligated lambda-gt 11 DNA was  
27 precipitated with 30 mM Na-acetate and 2.7 volumes of  
28 ethanol at -70°C for 2h. The lambda-gt 11 DNA ligated to  
29 cDNA was packaged in vitro at 20°C for 2 h using 25 µl of  
30 the lambda packaging mixture (Promega). The cDNA library  
31 was titrated on E. coli strain Y1090r on plates containing  
32 1mg/ml X-gal, and 0.4 mg/ml IPTG. The cDNA was amplified  
33 as plate lysate on E. coli strain Y1090r<sup>-</sup> at a density of  
15,000 plaques per 150mm plate.

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1

EXAMPLE 3

3 Screening the lambda-gt 11 cDNA library using specific  
monoclonal antibody probes.

5

Ryegrass pollen allergen-specific monoclonal  
7 antibodies were developed and characterized by SMART  
et al. (1983). Sera from patients allergic to ryegrass  
9 pollen were kindly provided by Dr David Hill from the  
Royal Children's Hospital, Melbourne.

11

The following procedure for screening the lambda-gt  
13 11 expression library is a modification of a previously  
described method (HUYNH, YOUNG AND DAVIS, 1985). A single  
15 colony of E. coli Y1090r<sup>-</sup> was grown at 37°C with good  
aeration to OD600 of 0.7 - 0.9, in LB medium containing  
17 100 ug/ml ampicillin and 0.4% (w/v) maltose. The cells  
were pelleted and resuspended in 10 mM MgSO<sub>4</sub> in 40% of  
19 the culture volume. The E. coli Y1090r<sup>-</sup> cells (0.3 ml)  
were then infected with approximately 18,000 - 20,000  
21 recombinant phage at 37°C for 15 minutes, plated onto  
150mB LB plates in 0.7% (w/v) agarose and incubated at  
23 42°C for 3 hours. The plates were overlaid with dry  
132mm nitrocellulose filters presoaked in 10mIPTG then  
25 incubated for 6 h at 37°C and the filters removed. A  
second IPTG-treated filter was placed on the bacterial  
27 lawn and the plates incubated overnight at 37°C. Filter  
plaque lifts were dried at room temperature, and washed  
29 with TBS for 10 minutes. The TBS was removed and 10ml of  
TBS containing 10% (w/v) non-fat milk powder was added and  
31 the filters were gently agitated for 1 h, drained, rinsed  
for 30 sec with TBS then washed for 10 minutes with TBS  
33 plus 0.1 (v/v) Tween -20 followed by two more washes of  
TBS for 10 min each. The filters were incubated for 3 h in

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1 TBS containing 2% (w/v) BSA and monoclonal antibodies  
2 (ascites) to ryetrass pollen allergens at a dilution of  
3 1:500 with gentle agitation. Following washing in  
4 TBS/0.1% (v/v) Tween-20, the filters were incubated ro 1.5  
5 h in TBS containing 2% (w/v) BSA and peroxidase-conjugated  
6 affinity - purified anti-mouse IgG at a dilution of  
7 1:500. The filters were washed and developed using fresh  
8 chromogenic peroxidase substrate 4-chloro-1-naphthol, 60mg  
9 dissolved in 20ml ice-cold methanol, and 80ml TBS  
10 containing 0.03% (v/v) H<sub>2</sub>O<sub>2</sub>. For each filter, 10ml of  
11 developing solution was used. After purple spots appeared  
12 on the filters, the developing solution was removed and  
13 the filters washed with distilled water to stop the  
14 reaction.

15

16 The developed filter was used to locate specific  
17 plaque areas on the plate, corresponding to a positive  
18 signal. Positive phage plaques were lifted from the  
19 plates a sagarose plugs and the eluted phage purified to  
20 individual antigen-positive lambda-gt 11 clones by  
21 rescreening at lower density on 85mm petri dishes with  
22 82mm nitrocellulose circles. Once plaque purification had  
23 been achieved, each of the lambda-gt 11 clones bearing  
24 allergen cDNA was lated at low denisty, and duplicate  
25 filter lifts were made. The ability of the recombinanat  
26 allergen to bind with antisera from allergic patients was  
27 detected using the same procudeure as described above,  
28 except that the overnight filter lift was incubated in the  
29 allergic antisera at a dilution of 1:10 whereas the first  
30 lift was treated with monoclonal antibodies.

31

#### EXAMPLE 4

32 Preparation of recombinant allergenic proteins from  
33 lambda-gt 11 recombinant lysogens and Western Blot  
34 Analysis.

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1

Bacteria (E. coli strain Y1089) to be lysogenized by the recombinant phage were grown to saturation in LB medium (pH 7.5) containing 0.2% (w.v) maltose at 37°C. One ml of the cells was collected by centrifugation and resuspended in 300 ul of LB medium containing 10mM MgCl<sub>2</sub>. The cells (approximately  $1 \times 10^8$  cells) were infected with about  $1 \times 10^9$  pfu of lambda-gt 11 recombinant phage containing cDNA inserts coding for the allergenic proteins (e.g. clone 12R and clone 6R) at 32°C for 20 minutes. The infected cells were serially diluted and plated at the density of 100-200 colonies per plate and incubated overnight at 32°C. Individual colonies were spotted onto replicate LB plates, of which one was incubated at 42°C, and the other at 32°C overnight. Recombinant lysogen clones were indicated by growth at 32°C but not at 42°C, and occurred at a frequency of 20% for clone 12R, and 3-4% for clone 6R.

19

In order to obtain a preparative amount of the recombinant allergenic proteins, a single lysogen colony of Y1089 was inoculated into 10ml of LB medkium and incubated 32°C with good aeration until the OD600 reached 0.5. The culture was quickly shifted to a 42°C water bath and incubated for 20 minutes with shaking.

27

The lac operon repressor was inactivated by addition of 100 ul of 1M IPTG. The culture was then incubated at 37°C for 1 hour allowing the lac Z gene to be expressed and the allergenic proteins to be synthesized as a fusion protein with beta-galactosidase. Cells were harvested by spinning at 3000 rpm for 10 minutes at room temperature, resuspended in 150 ul of Sample buffer and immediately frozen in liquid nitrogen. The cells were

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1 lysed by thawing at room temperature. For electrophoretic  
analysis of proteins, 150 ul of SDS sample buffer  
3 containing bromophenol blue tracking dye was added to the  
freeze-thaw lysate. Samples were boiled for 3 minutes and  
5 the insoluble material removed by micro-entrifugation for  
3 minutes.

7  
Proteins were resolved by 7-10% (w/v) SDS -  
9 polyacrylamide gel electrophoresis and visualised by  
Coomassie Blue-staining with duplicate samples  
11 electroblotted onto nitrocellulose filter using the  
Bio-Rad Trans Blot apparatus (0.15 amps overnight).  
13 Fusion proteins were detected with monoclonal antibodies  
and visualized using the screening procedure described  
15 previously.

17 EXAMPLE 5

Northern analysis.

19  
Total RNA was extracted from pollen, leaf, hydrated  
21 seed and root samples as previously described for pollen,  
and 20 ug RNA/sample electrophoresed in formaldehyde/1.2%  
23 (w/v) agarose gels (MANIATIS et al., 1982) run at 70V for  
4 hrs in running buffer containing 20mM morpholinopropane  
25 sulphonic acid, 5mM sodium acetate and 0.1mM EDTA, to pH  
7.0. The RNA's were transferred to nitrocellulose (Hybond  
27 C) filters and pre-equilibrated 2 hours at 50°C in  
hybridization buffer containing 50% (v/v) deionised  
29 formamide, 2X SSPE, 7 % (w/v) SDS, 0.5% (w/v) non-fat milk  
powder, 1% PEG 20,000, and 0.5mg/ml non-homologous herring  
31 sperm carrier DNA. Fresh hybridization buffer containing  
the random primed Lol pI DNA probe was added and incubated  
33 at 50°C for overnight hybridization. Filters were washed  
vigorously in 2X SSC, 0.1% (w/v) SDS for 15 minutes at RT,

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1 then 0.5 X SSC, 1% (w/v) SDS at 50°C for 15 minutes,  
followed by a brief rinse in 0.5x SSC, 0.1% (w/v) SDS,  
3 blotted lightly and wrapped in Glad Wrap. Kodak film was  
exposed for 18 hours at -70°C.

5

EXAMPLE 6

7 Expression of Lol pI cDNA products reacted with IgE from  
allergic sera.

9

The cDNA insert from lambda-gt 11-12R which codes  
11 for Lol pI was sub-cloned into the EcoRI site of the  
plasmid expression vector pGEX where it can be expressed  
13 as a fusion protein with glutathione transferase. E. coli  
infected with this plasmid pGEX-12R or with the  
15 non-recombinant vector alone, were grown at a log phase  
culture, and the bacteria pelleted by centrifugation.  
17 These bacteria were lysed and the total proteins separated  
on SDS-PAGE gel. A western blot shows that only bacteria  
19 containing recombinant-plasmids possess a protein  
component reactive with specific IgE in sera taken from  
21 donors known to be allergic to ryegrass pollen. Those  
results are shown in Figure 6.

23

EXAMPLE 7

25 Cross-reactivity of Lol pI with homologous allergens from  
other grass pollen.

27

Lol pI is a protein of MW 34 kD, and SDS-PAGE shows  
29 that other common grasses possess a homologous protein of  
similar molecular weight. Our results show that these  
31 proteins share a common antigenic epitope (detected by  
monoclonal antibodies), and are allergens in terms of  
33 specific IgE- binding. Results are shown in Figure 7.  
Because of this allergenic similarity, Lol pI is the



- 51 -

1 immunodominant allergen of grass pollen. A consequence is  
that the cDNA clone 12R can be used as a heterologous  
3 probe to isolate the homologous cDNA clones for allergens  
from other grass pollens.

5

- 52 -

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27

## CLAIMS

1. A recombinant vector comprising a DNA sequence encoding a protein displaying allergenic activity from pollen of a grass species.
2. The recombinant vector according to claim 1, wherein the allergenic protein is from pollen of grass belonging to the family Poaceae (Gramineae).
3. The recombinant vector according to claim 2, wherein the allergenic protein is from pollen of grass belonging to the genus Lolium.
4. The recombinant vector according to claim 3, wherein the allergenic protein is immunologically cross-reactive with antibody to Lol pI protein of Lolium perenne pollen, namely:

Pooid (festucoid) grasses. Group 1: Triticanae: Bromus inermis, smooth brome; Agropyron repens, English couch; A. cristatum; Secale cereale, rye; Triticum aestivum, wheat. Group2: Poanae: Dactylis glomerata, orchard grass or cocksfoot; Festuca elatior, meadow fescue; Lolium perenne, perennial ryegrass; L. multiflorum, Italian ryegrass; Poa pratensis, Kentucky bluegrass; P. compressa, flattened meadow grass; Avena sativa, oat; Holcus lanatus, velvet grass or Yorkshire fog; Anthoxanthum odoratum, sweet vernal grass; Arrhenatherum elatius, oat grass; Agrostis alba, red top; Phleum pratense, timothy; Phalaris arundinacea, reed canary grass.

Panicoid grass, Paspalum notatum, Bahia grass,  
Andropogonoid grasses: Sorghum halepensis, Johnson grass;  
Zea mays, maize.

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5. The recombinant vector according to claim 4 wherein the allergenic protein is Lol pI of ryegrass, Lolium perenne, pollen, or a derivative or homologue thereof.
6. The recombinant vector according to claim 5 comprising a DNA sequence as depicted in Figure 5 or a degenerate or homologous form thereof.
7. A recombinant DNA molecule comprising a eukaryotic or prokaryotic origin of replication, a detectable marker, a DNA sequence encoding the Lol pI allergenic protein or a derivative or a homologue thereof or an allergenic protein cross-reactive with an antibody to said Lol pI protein or its derivatives or homologues and optionally a promoter sequence capable of directing transcription of said DNA sequence.
8. The recombinant DNA molecule according to claim 7 comprising a DNA sequence as depicted in Figure 5 or a degenerate or homologous form thereof.
9. The recombinant DNA molecule according to claim 7 or 8, wherein the promoter is the Lol pI gene promoter.
10. A host cell carrying a vector or recombinant DNA molecule according to anyone of claims 1 to 9.
11. A method for isolating and identifying DNA encoding an allergenic protein of pollen from the family Poaceae (Gramineae) comprising screening by hybridization DNA isolated from said family with a DNA or RNA sequence encoding Lol pI protein of Lolium perenne pollen or its derivatives or homologues .

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12. The method according to claim 11, wherein the DNA to be identified comprises DNA from a cDNA library, which is prepared by reverse transcription on a template of mRNA of grass pollen showing allergenic activity.
13. A method of producing recombinant Lol pI or a derivative or homologue thereof or an allergenic protein immunologically reactive to antibodies to Lol pI or a derivative or homologue thereof, comprising culturing an organism containing a replicable recombinant DNA molecule, said molecule comprising a promoter capable of expression in said organism, the gene encoding Lol pI or its derivative or homologue or an immunologically related protein of Lol pI located downstream of and transcribed from said promoter, a selectable marker and a DNA vehicle containing a prokaryotic or eukaryotic origin of replication, under conditions and for a time sufficient for said recombinant DNA molecule to be stably maintained and direct the synthesis of Lol pI or its derivative, homologue or immunological relative and then isolating same.
14. The method according to claim 15, wherein the promoter is the Lol pI promoter or homologue or degenerate form thereof and the host organism is one in which said promoter will function.
15. Non-native Lol pI or a derivative or homologue thereof or a non-native allergenic protein immunologically reactive to antibodies to said Lol pI or its derivative or homologue.



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16. An antibody to non-native Lol pI or a derivative or homologue thereof or to a non-native allergenic protein immunologically reactive to antibodies to said Lol pI or its derivative or homologue.

17. A method of detecting Lol pI or a derivative or homologue thereof or an allergenic protein immunologically reactive with said Lol pI or its derivative or homologue in serum, tissue extract, plant extract or other biological fluid comprising the steps of contacting said serum, extract or fluid to be tested with an antibody according to claim 14 for a time and under conditions sufficient for an allergenic protein-antibody complex to form and subjecting said complex to a detecting means.

18. The method according to claim 17, wherein the antibody is labelled with a material providing a detectable signal, said material selected from the group consisting of a radioactive isotope, and enzyme, a fluorescent molecule, a chemiluminescent molecule, a bioluminescent molecule or a cell.

19. The method according to claim 17, wherein the allergenic protein-antibody complex is detected by contacting said complex with a second antibody specific to the first antibody, said second antibody labelled with a material providing a detectable signal, said material selected from the group consisting of a radioactive isotope, an enzyme, a fluorescent molecule, a chemiluminescent molecule, a bioluminescent molecule or a cell for a time and under conditions sufficient for a tertiary complex to form and then detecting said signal.

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20. A method of detecting an antibody to an allergenic protein from pollen of the family Poaceae (Gramineae) in serum or other biological fluid comprising contacting said serum or fluid with recombinant Lol pI or its antigenic derivative for a time and under conditions sufficient for an antibody - Lol pI complex to form and subjecting said complex to a detecting means.
21. The method according to claim 20, wherein the recombinant Lol pI or its antigenic derivative is optionally labelled with a reporter molecule.
22. The method according to claim 20, wherein the complex is detected by contacting said complex with a second antibody specific to said Lol pI or its antigenic derivative and said second antibody being labelled with a reporter molecule, for a time and under conditions sufficient for a tertiary complex to form and then detecting said reporter molecule.
23. The method according to claim 21 or 22 wherein said reporter molecule is selected from the group consisting of a radioactive isotope, and enzyme, a fluorescent molecule, a chemiluminescent molecule a bioluminescent molecule or a cell.
24. A kit for the detection of antibodies to a protein having allergenic properties, said protein from pollen of the family Poaceae (Gramineae), the kit being compartmentalized to receive a first container adapted to contain recombinant Lol pI or its antigenic derivative or homologue, and a second container adapted to contain an antibody to Lol pI or its derivative or homologue, said antibody labelled with a reporter molecule capable of giving a detectable signal.

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25. The kit according to claim 24, wherein the reporter molecule is a radioisotope, an enzyme, a fluorescent molecule, a chemiluminescent molecule, a bioluminescent molecule or a cell.

26. The kit according to claim 25, wherein the reporter molecule is an enzyme.

27. The kit according to claim 26, wherein the kit further comprises a third container adapted to contain a substrate for the enzyme.

28. The kit according to claim 24 alternatively comprising a container adapted to contain recombinant Lol pI or is antigenic derivative or homologue labelled with a reporter molecule capable of giving a detectable signal.

29. A recombinant DNA molecule comprising a ryegrass pollen promoter sequence or homologue or degenerate form thereof located on said molecule and further having one or more restriction endonuclease sites downstream of said promoter such that a nucleotide sequence inserted into one or more of these sites is transcribeable in the correct reading frame.

30. The recombinant DNA molecule according to claim 29, wherein said promoter is the Lol pI gene promoter.

31. The recombinant DNA molecule according to claim 30 further comprising a selectable marker.

32. The recombinant molecule according to claim 29 or 30 or 31 further comprising means for stable inheritance in a prokaryotic and/or eukaryotic cell.

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33. The recombinant DNA molecule according to claim 32, wherein said means comprises a prokaryotic or eukaryotic origin of replication thereby permitting said molecule to replicate extrachromosomally in a host cell.

34. The recombinant DNA molecule according to claim 33, wherein said molecule is replicable in prokaryotic cells.

35. The recombinant DNA molecule according to claim 34, wherein the prokaryotic cells comprise Escherichia coli, Pseudomonas or Bacillus.

36. The recombinant DNA molecule according to claim 33, wherein said molecule is replicable in eukaryotic cells.

37. The recombinant DNA molecule according to claim 36, wherein the eukaryotic cells comprise cells from yeast, insects, mammals or plants.

38. The recombinant DNA molecule according to claim 37, wherein the eukaryotic cells are plant cells derived from the family Poaceae.

39. The recombinant DNA molecule according to claim 32, wherein said molecule replicates by insertion into the genome of a host cell and replicates in synchrony with said genome.

40. The recombinant DNA molecule according to any one of claims 29 to 39 further comprising a nucleotide sequence encoding a polypeptide or portion thereof or a mRNA or a portion thereof inserted into one of the restriction endonuclease sites downstream of said promoter such that said nucleotide sequence is transcribeable in the correct reading frame.

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41. The recombinant DNA molecule according to claim 40, wherein said nucleotide sequence encodes an allergenic protein, a cytokinin or a protein having a deleterious function on a plant cell, or their derivatives.

42. The recombinant DNA molecule according to claim 41, wherein the allergenic protein in Lol pI or its derivative.

43. The recombinant DNA molecule according to claim 41, wherein the nucleotide sequence encodes a toxin, said toxin active against cells derived from the family Poaceae.

44. The recombinant DNA molecule according to claim 40, wherein the nucleotide sequence encodes an antisense RNA capable of inhibiting translation of a gene in a cell from the family Poaceae.

45. A prokaryote or eukaryote transformed with a recombinant DNA molecule according to any one of the preceding claims.

46. A method of inhibiting pollen development or function and thereby inducing nuclear male sterility in plants of the family Poaceae comprising the steps of:

a) developing a plant carrying a recombinant DNA molecule comprising the rye grass pollen promoter sequence or homologue or degenerate form thereof located on said molecule and a nucleotide sequence encoding a polypeptide having a deleterious function in cells derived from the family Poaceae, said nucleotide sequence transcribeable from said promoter, and said recombinant DNA molecule stably contained in pollen producing cells, and,

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b) growing said plants under conditions and for a time sufficient for their developmental stage to cause expression of said nucleotide sequence from said promoter thereby producing the polypeptide having a deleterious function on said pollen producing cells such that pollen formation is inhibited or said pollen is inactive.

47. The method according to claim 46, wherein the nucleotide sequence alternatively encodes an antisense RNA capable of inhibiting pollen formation of rendering said pollen inactive.

48. A method for desensitizing a human allergic to a grass pollen comprising administering to said human a desensitizing effective amount of Lol pI or a derivative, homologue or immunological relative thereof for a time and under conditions sufficient to effect desensitization of said human.

49. The method according to claim 48 wherein administration is by the intravenous, intramuscular, intranasal, intradermal, intraperitoneal, suppository or oral route.

50. A pharmaceutical composition useful in desensitizing a human allergic to a grass pollen comprising an effective amount of Lol pI, or a derivative, homologue or immunological relative thereof, and a pharmaceutically acceptable carrier.

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51. A peptide, polypeptide or protein comprising an amino acid sequence corresponding in whole or part to the nucleotide coding sequence represented in Figure 5 or to degenerate or homologue forms thereof.

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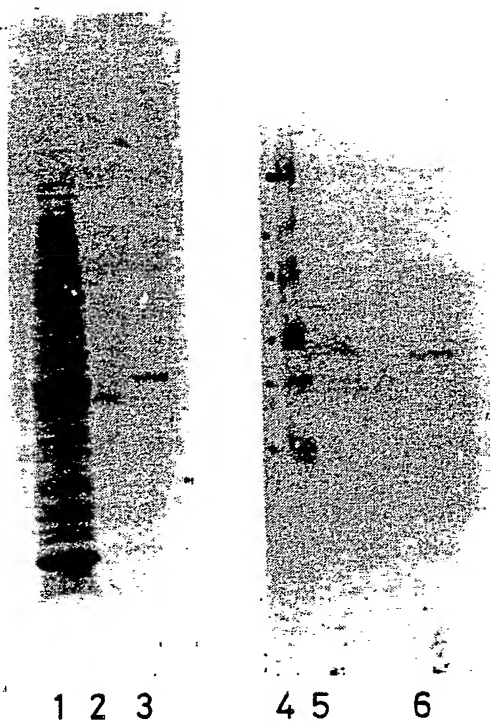


Fig.1.

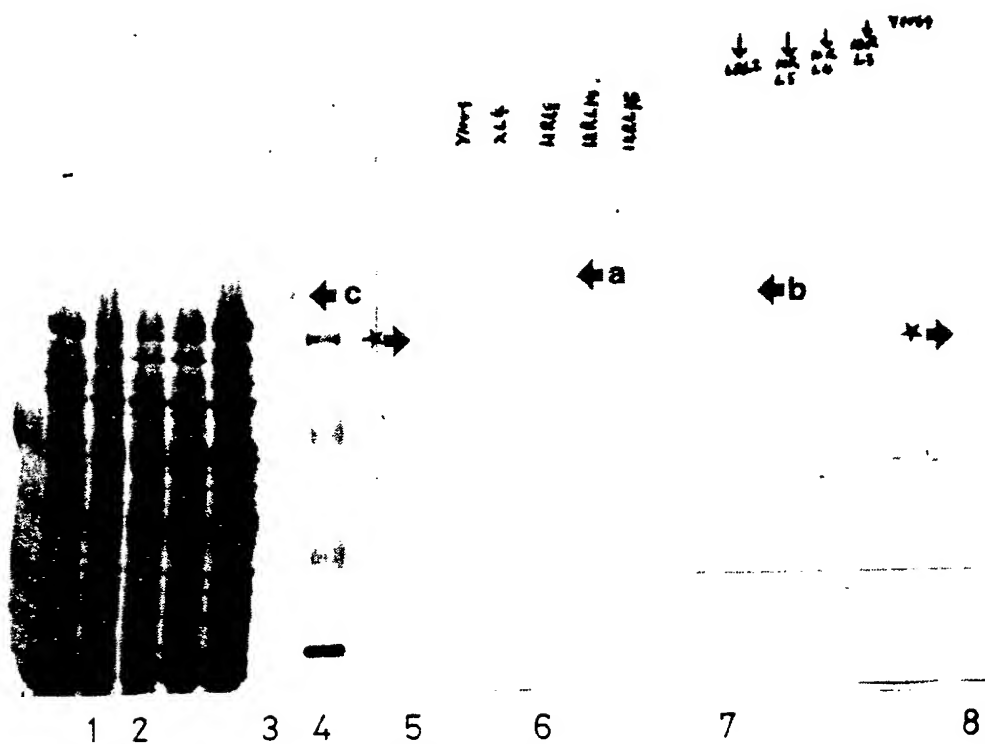


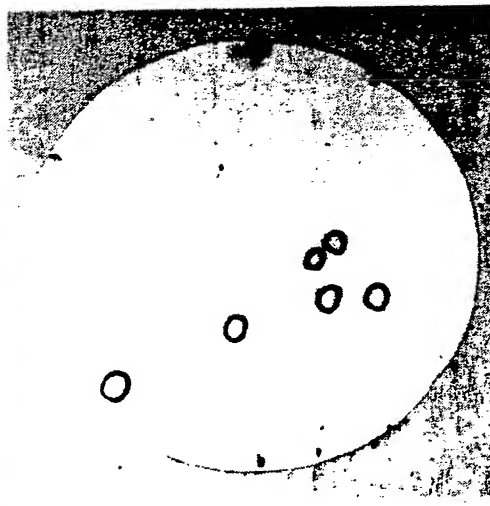
Fig.3.



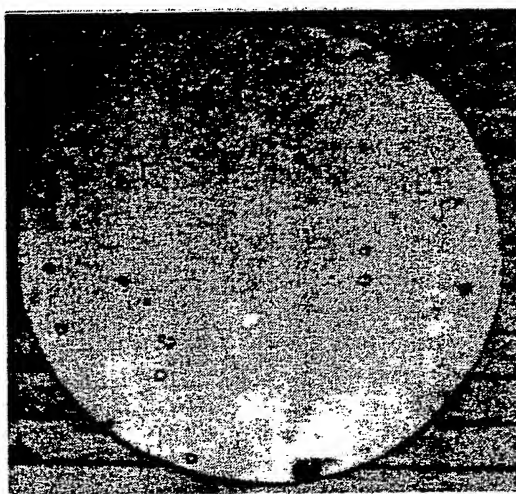
2/12



*Fig. 2a.*



*Fig. 2c.*



*Fig. 2b.*



*Fig. 2d*

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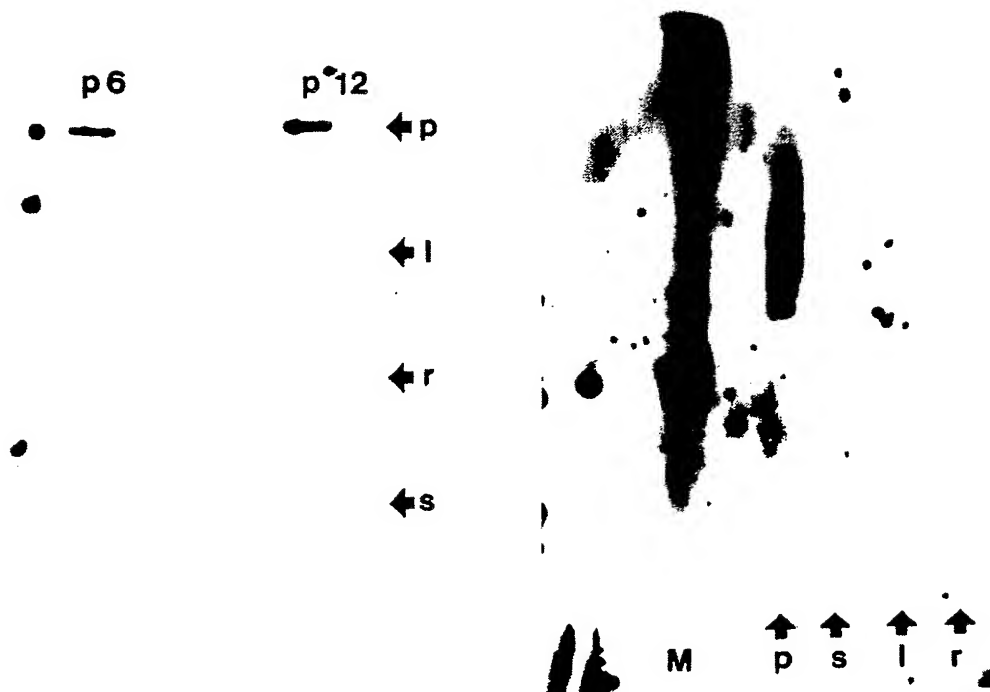


Fig.4a.

Fig.4b.

1 2 3 4

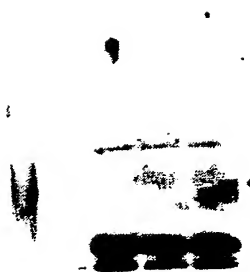


Fig.6.

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10	20	30	40	50	60
GAATTCCGCT	ATCCCTCCCT	CGTACAAACA	AACGCAAGAG	CAGCAATGGC	CGTCCAGAAC
70	80	90	100	110	120
TACACGGTGG	CTCTATTCTT	CGCCGTGGCC	CTCGTGGGGG	CCCGGCCGCT	CCTACGCCGC
130	140	150	160	170	180
TGACGCCGGC	TACACCCCCG	CAGCCGGGCC	ACCCCGGCTA	CTCCTGCTGC	CACCCCGGCT
190	200	210	220	230	240
GCGGCTGGAG	GGAAGGCGAC	GACCGACGAG	CAGAAGCTGC	TGGAGGACGT	CAACGCTGGC
250	260	270	280	290	300
TTCAAGGCAG	CCGTGGCCGC	CGTGCCCAACG	CCCCTCCGGC	GGACAAGTTC	AAGATCTTCG
310	320	330	340	350	360
AGGCCGCCCTT	CTCCGAGTCC	TCCAAGGGCC	TCCTCGCCAC	CTCCGCCGCA	AGGCACCCCGG
370	380	390	400	410	420
CCTCATCCCC	AAGCTCGACA	CCGCCCTACGA	CGTCGCTACA	AGGGCGAGGG	CGCCGCCACC

Fig.5(1).

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430	440	450	460	470	480
CCCGAGGCCA	AGTACGACGC	CTTCGTCACT	GCCCTCACCG	AAGCTCCGCG	TCATCGCCGG
490	500	510	520	530	540
CGCCCTCGAG	GTCCACGCCG	TCAAGCCCCG	CACCGAGGAG	GTCCCTGCTG	CTAAGATCCC
550	560	570	580	590	600
CACCGGTGAG	CTGCAGATCG	TTGACAAGAT	CGATGCTGCC	TTCAAGATCG	CAGCCACCGC
610	620	630	640	650	660
CGCCGCCAAC	GCCGCCCCCA	CCAACGATAA	GTTACCCGTC	TTGAGAGGTG	CCTTCAACAA
670	680	690	700	710	720
GGCCCTCAAT	GAGTGCACCG	GGCGGCGCTA	TGAGACCTAC	AAGTTTATCC	CCTCCCCTCGA
730	740	750	760	770	780
GGCCGCGGTC	AAGCAGCCCTA	CGCCGCCACC	GTCCGCCGCG	GCCCGAGGTC	AAGTACCGCC

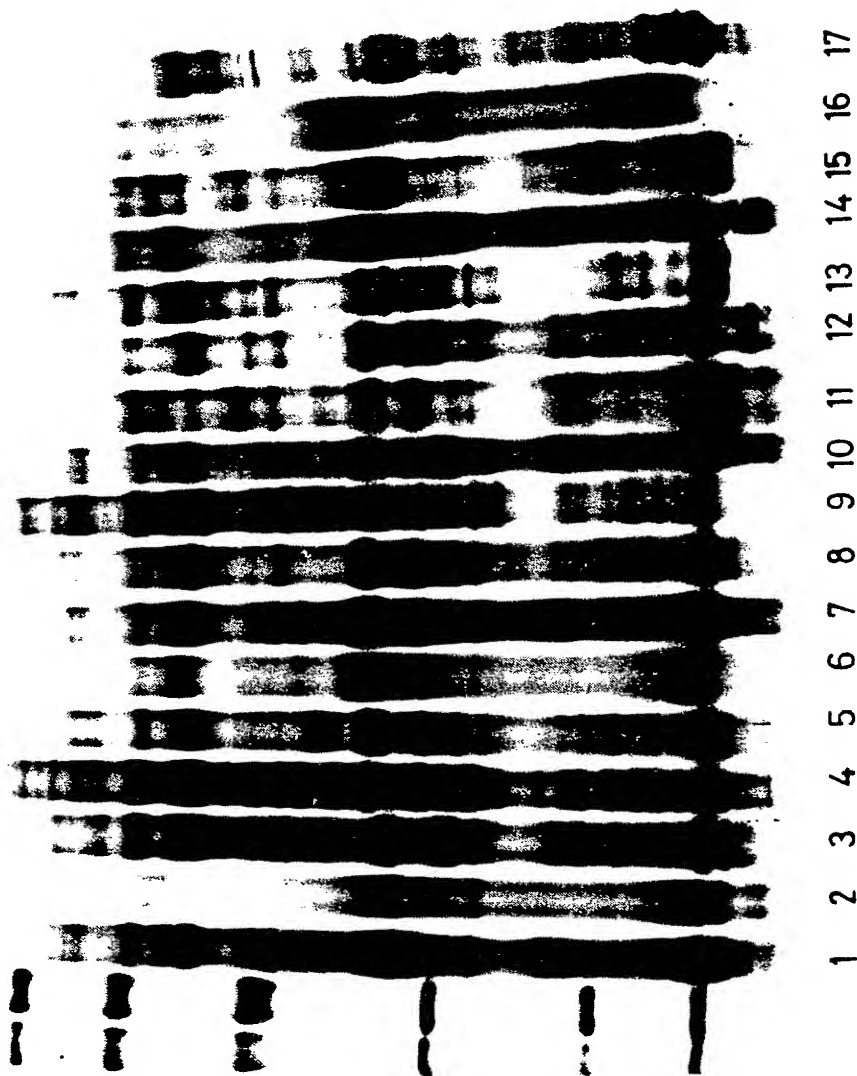
Fig.5(2).

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790	800	810	820	830	840
GTCTTTGAGG	CCGCGCTCGA	CCAAGGCCAT	CACCGCCATG	ACCCAGGCAC	AGAAGGCCGG
850	860	870	880	890	900
CAAACCCGCT	GCCGCCGCTG	CCACAGGCCG	CAACCGTTGC	CACCGCACCG	CAACCGCCGC
910	920	930	940	950	960
C--TG-C-CA	G-CCGCCGCT	GCTGCTGGCT	ACCAAAGCCT	GATCAGCTTG	CTAATATACT
970	980	990	1000	1010	1020
ACTGAACGTA	TGTATGTGCA	TGATCCGGGC	GGCGAGTGGT	TTTGTTGATA	ATTAATCTTC
1030	1040	1050	1060	1070	1080
GTTTTCGTTT	CATGCAGCCG	CGATCGAGAG	GTTCGATCGT	TGTAATAATT	CAATATTTT
1090	1100	1110	1120	1130	1140
TATTTCTTTT	TGAATCTGTA	AATCCCCATT	GACAAGTAGT	GGGATCAAGT	CG-CATGTAT
1150	1160	1170	1180	1190	1200
CACCGTTGAT	GCTGAGTTTA	ACGATGGGGA	GTTTATCAAA	GAATTTATTA	TTAAAAAAA
1210	1220	1230	1240	1250	1260
AAAAAAAAAA	AAAAAAAAAA	AAAAAAAAAA	AAAAAAAAAA	AAAAA	

*Fig.5(3).*

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*Fig. 7a.*

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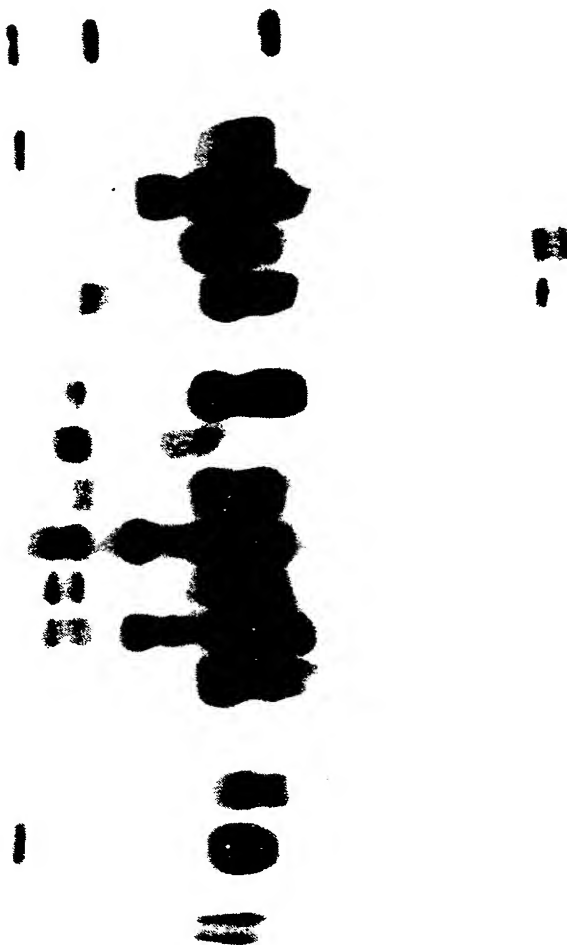
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17



*Fig. 7b.*

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1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17



Lot p1 ➤

Fig.7c.



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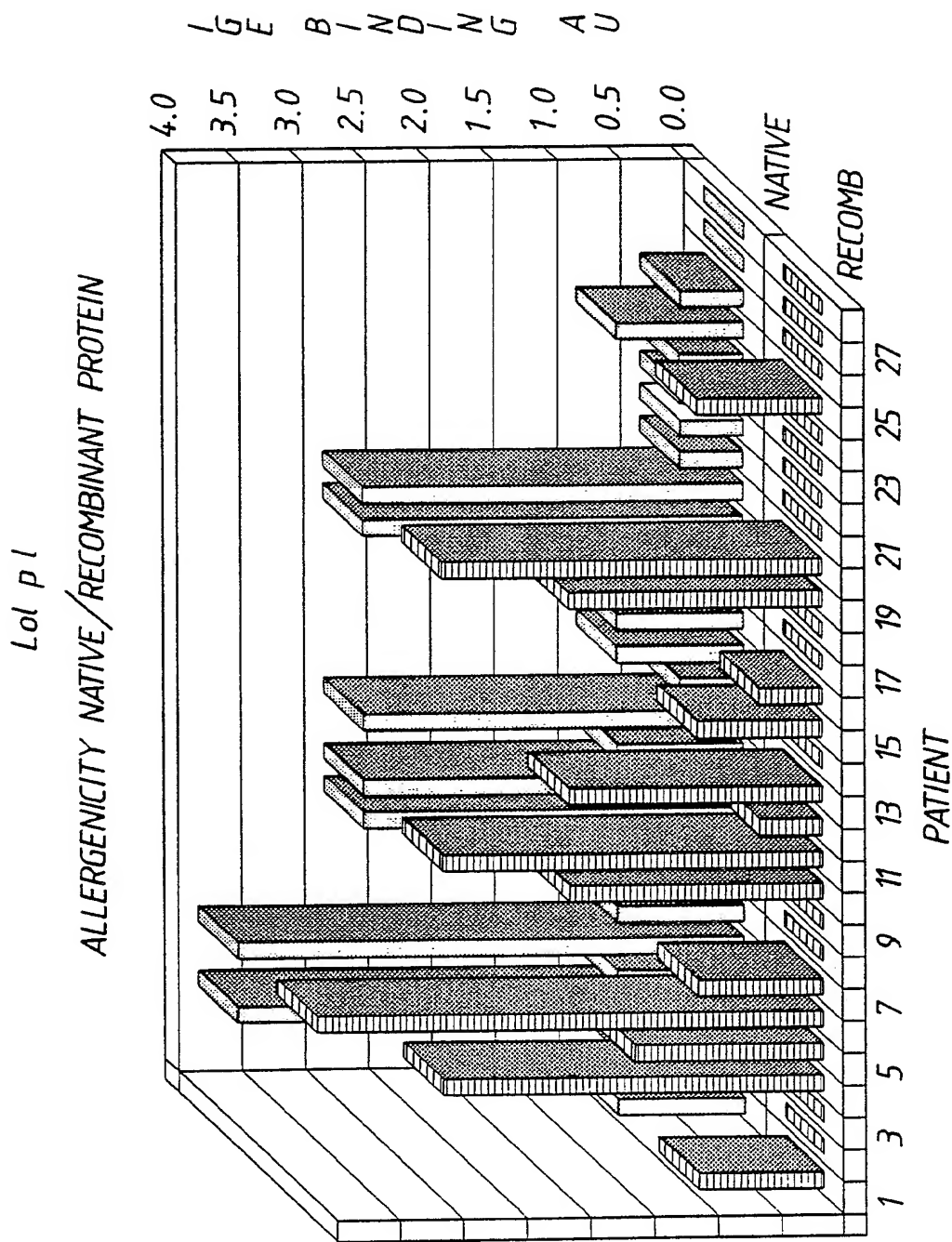


Fig. 8a.

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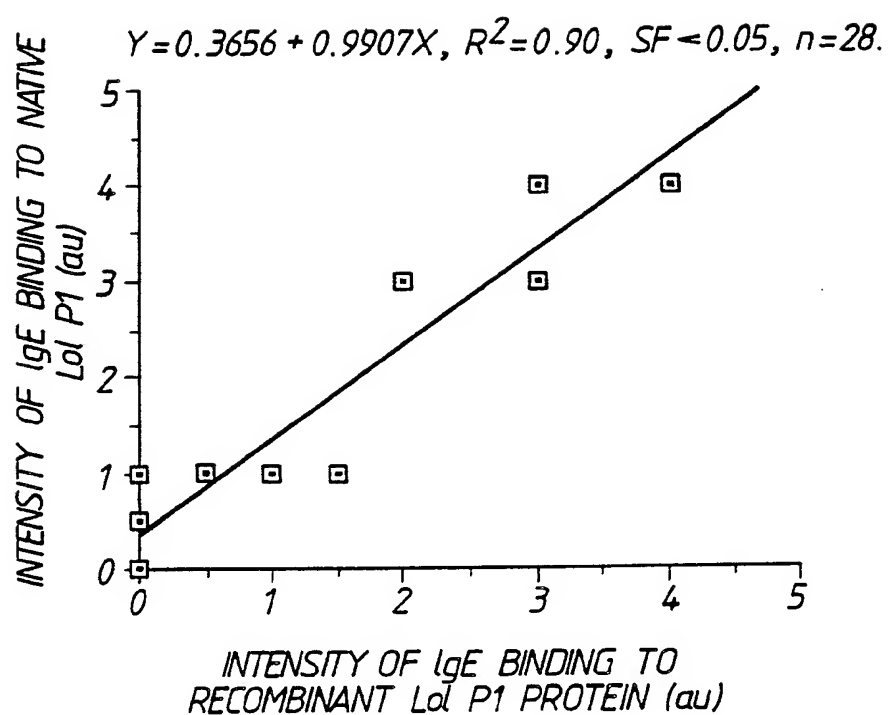


Fig.8b.

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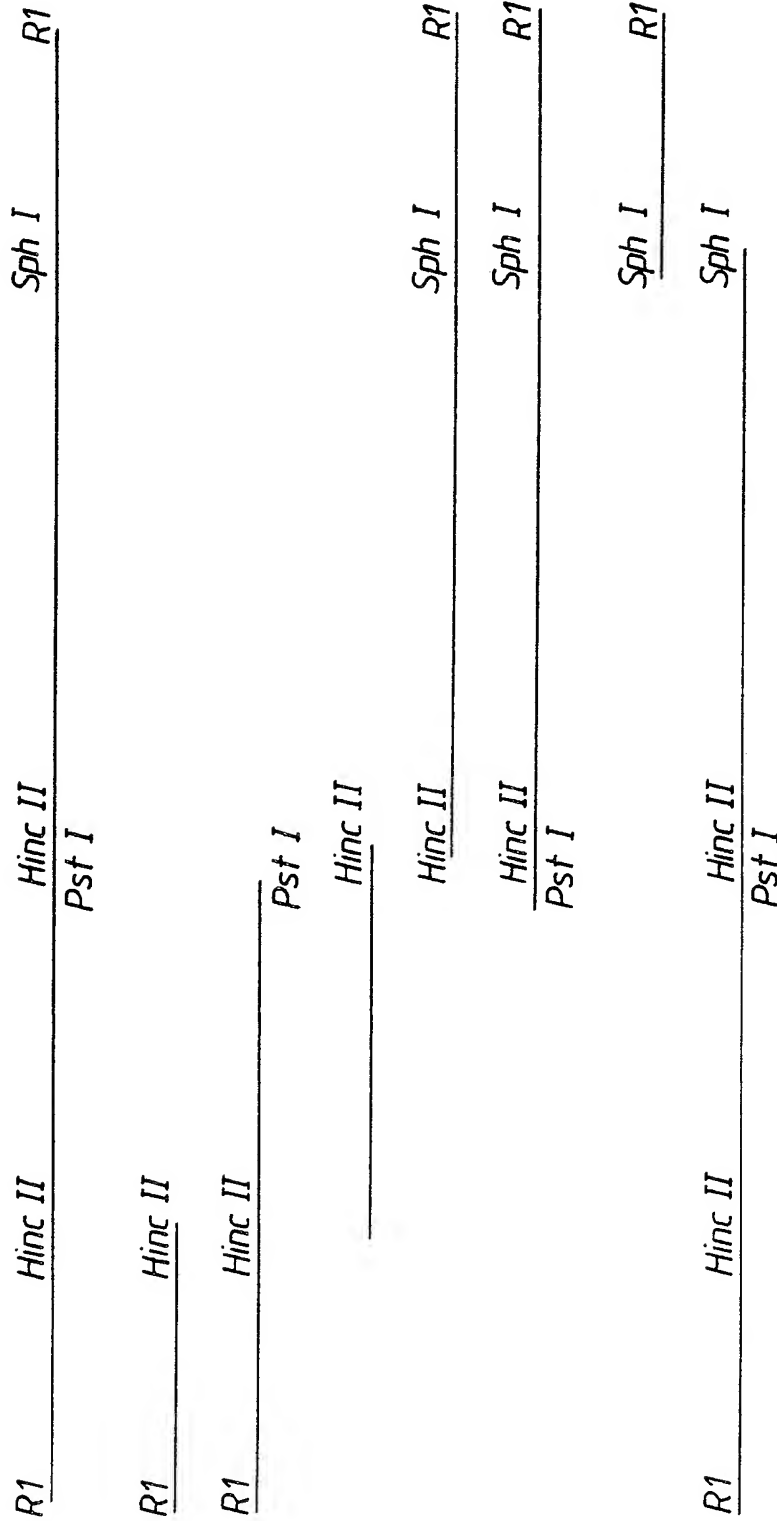


Fig.9.

## I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) 6

According to International Patent Classification (IPC) or to both National Classification and IPC  
 Int. Cl.<sup>4</sup> C12N 15/00, 1/20, 5/02, C12P 21/02, 19/34, C07K 13/00, G01N 33/531, 33/532,  
 C07K 15/12, 15/14, C12Q 1/68, A01H 1/00, C07H 21/04, A61K 39/36

## II. FIELDS SEARCHED

Minimum Documentation Searched 7

Classification System |

Classification Symbols

IPC

WP1, WP1L, USPA, DERWENT DATA BASES: KEYWORDS RYE GRASS POLLEN ALLERGEN  
 OR ANTIGEN; RECOMBINANT L01 PI

Documentation Searched other than Minimum Documentation  
 to the Extent that such Documents are Included in the Fields Searched 8

AU : C12N 15/00, C07K 15/12, 15/14,  
 CHEMICAL ABSTRACTS, BIOSIS PREVIEWS, EXERPTA MEDICA, MEDLINE KEYWORDS as above

## III. DOCUMENTS CONSIDERED TO BE RELEVANT 9

Category*	Citation of Document, with indication, where appropriate, of the relevant passages 12	Relevant to Claim No 13
X	Molecular Immunology, volume 23, no. 12, 1986 pp. 1281-1288, C.R. Kahn and D.G. Marsh, "Monoclonal Antibodies to the major <u>Lolium perenne</u> (Rye Grass) pollen allergen L01 pI (Rye I)".	(16-18, 20-23, 24-25, 28, 51)
X	Immunology, vol 59 no. 2, 1986 pp. 309-315, R. Bose et al., "Production and characterization of mouse mono clonal antibodies to allergenic epitopes on L01pI (Rye I)".	(16-17, 20, 21, 23, 51)
Y		(24-28)
X	Int. Arch. Allergy, Appl. Immun. vol. 72 no. 3 pp. 243-248, 1983, I.J. Smart et al., "Development of Monoclonal mouse antibodies specific for allergenic components in Ryegrass ( <u>Lolium perenne</u> ) pollen".	(16, 17, 19, 24-28, 51)
X	Int. Arch. Allergy, Appl. Immun. vol. 78, 1985 pp. 300-304, M.B. Singh & R.B. Knox, Grass Pollen Allergens: "Antigenic Relations ships Detected Using monoclonal antibodies and Dot blotting Immunoassay".	(16-17, 19)
Y	(CONTINUED)	(24-28)
* Special categories of cited documents: 10		
"A"	document defining the general state of the art which is not considered to be of particular relevance	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"E"	earlier document but published on or after the international filing date	"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step
"L"	document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.
"O"	document referring to an oral disclosure, use, exhibition or other means	"&" document member of the same patent family
"P"	document published prior to the international filing date but later than the priority date claimed	

## IV. CERTIFICATION

Date of the Actual Completion of the  
 International Search  
 30 June 1989 (30.06.89)

Date of Mailing of this International  
 Search Report

11 July 1989 (11.07.89)

International Searching Authority

Signature of Authorized Officer

Australian Patent Office

J H CHAN



## FURTHER INFORMATION CONTINUED FROM THE SECOND SHEET

X	Journal of Allergy and Clinical Immunology vol 78 no 6 pp. 1190-1201, 1986 L.R. Friedhoff et al., "A study of the human immune response to <u>Lolium perenne</u> (Rye) pollen and its components, L01 pI and L01 pII (Rye I and Rye II)".	(16, 20-21, 23, 24-28, 51)
X,P	Tissue Antigens vol 31 no 4 pp 211-219 (1988) L.R. Friedhoff et al., "Association of HLA-DR3 with human immune response to L01 pI and L01 pII allergens in allergic subjects".	(20-21, 23-28, 50, 51)
X	Int. Arch. Allergy, Appl. Immun. volume 85 no.1 pp 104-108 (1988) R.B. Cook et al "Induction of Allergen -Specific T-Cells by conjugates of N-formyl-methionyl-leucyl-phenylalanine and Rye grass pollen extract".	(48-51)
Y,P	Chemical abstracts vol. 108 issue no 23 1988. Wheeler A.W. et al "Retained T-cell reactivity of rye grass pollen extract following cleavage with cyanogen bromide and nitrothiocyanobenzoic acid". (CONTINUED)	(48-51)

## V. [X] OBSERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 1

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. [X] Claim numbers ...48-50, because they relate to subject matter not required to be searched by this Authority, namely:  
Rule 39(iv) Methods for treatment of the human or animal body by surgery or therapy.
2. [] Claim numbers ..., because they relate to parts of the international application that do comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. [] Claim numbers ..., because they are dependent claims and are not drafted in accordance with the second and third sentences of PCT Rule 6.4 (a):

## VI. [X] OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING 2

This International Searching Authority found multiple inventions in this international application as follows:

- Group I - Claims 1-14, 20-28 and 42 are directed to recombinant known L01 pI protein, cDNA thereof, expression of cDNA in transformed host, cDNA as probe, use of recombinant protein.

(CONTINUED)

1. [] As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims of the international application.
2. [] As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims of the international application for which fees were paid, specifically claims:
3. [X] No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claim numbers:  
1-14, 20-28, 42, 15-19, 48-50 and 51.
4. [] As all searchable claims could be searched without effort justifying an additional fee, the International Searching Authority did not invite payment of any additional fee.

## Remark on Protest

- [ ] The additional search fees were accompanied by applicant's protest.  
[ ] No protest accompanied the payment of additional search fees.

III. DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)		
Category*	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim No
X	Journal of Chromatography, vol 370 issue no. 1 pp. 165-172 (1986) BRIEVA A. and RUBIO N. "Rapid purification of The Main allergen of <u>Lolium perenne</u> by high performance liquid chromatography".	(51)